Paroxysmal nocturnal haemoglobinuria: Nature’s gene therapy?

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The development of paroxysmal nocturnal haemoglobinuria (PNH) requires two coincident factors: somatic mutation of the PIG-A gene in one or more haemopoietic stem cells and an abnormal, hypoplastic bone marrow environment. When both of these conditions are met, the fledgling PNH clone may flourish. This review will discuss the pathophysiology of this disease, which has recently been elucidated in some detail.

The development of paroxysmal nocturnal haemoglobinuria (PNH) requires two coincident factors: somatic mutation of the PIG-A gene in one or more haemopoietic stem cells and an abnormal, hypoplastic bone marrow environment. When both of these conditions are met, the PNH clone may flourish. Recently, the pathophysiology of this disease has been elucidated in some detail and we now have rational theories concerning the clinical manifestations of PNH, its association with other haematological disorders (such as aplasia and myelodysplasia), and the sequence of events that leads to overt disease. Spontaneous somatic mutations are common in the PIG-A gene, leading to failure of synthesis of the glycosyl phosphatidylinositol (GPI) anchor. Without this structure, many proteins are unable to attach to cell surfaces. Red blood cells lose complement defence proteins, which explains the classic feature of intravascular complement mediated haemolysis. There is indirect evidence that platelet activation with consequent thrombosis is caused by a similar mechanism. The relative growth advantage of PNH cells in a hypoplastic marrow is also, presumably, a direct or indirect result of these alterations in surface antigen composition, although the precise pathophysiological mechanisms remain to be described. It is probable that the association with aplasia is explained by this relative growth advantage and that clonal evolution to a leukemic state is a consequence of the primary insult causing the aplasia. In this way, PNH can be seen as an attempt to restore a form of useful, if abnormal, haemopoiesis in a damaged bone marrow: nature’s gene therapy.

THE EVOLUTION OF OUR UNDERSTANDING OF PNH
PNH as a clinical entity has puzzled physicians and scientists for 200 years. Perhaps the first description was “An account of a singular periodic discharge of blood from the urethra” written in 1794 by a Scottish surgeon, Charles Stewart. However, the first detailed description is credited to Paul Strübing in 1882. His patient had a six year history of passing dark urine intermittently in the mornings, always clearing by noon. His conclusions were detailed and astonishingly perceptive, suggesting as he did that there was intravascular haemolysis and that some of the patient’s symptoms were the result of thrombosis. He even concluded (correctly) that a red blood cell defect was to blame. Despite this insight, the work was largely ignored and when Marchiafava reported a case in Italy 29 years later it was regarded as a new entity. Marchiafava–Michelli published further observations in 1931 and used the weighty term “spleenomegalic haemolytic anaemia with haemoglobinuria” and haemosiderinuria, Marchiafava–Michelli type” from which the disease retains the eponym. The modern term PNH was first coined shortly before Michelli’s paper in a description of a case from the Netherlands.

Attempts to explain the haemolysis began in Rotterdam in 1911, when Hymans van den Bergh noted that PNH erythrocytes were sensitive to lysis in vitro when exposed to carbon dioxide. By the 1930s it had been shown that the lysis was complement mediated and pH dependent: being greatest in acidified conditions. This formed the basis for Ham’s test, which was the diagnostic gold standard until its replacement by flow cytometry in recent years.

In 1944, Sir John Dacie first noted the association of PNH with aplasia in a case of Fanconi’s anaemia. In the 1950s, the defect was shown to be present in other haemopoietic lineages, with the observation that the neutrophil alkaline phosphatase was reduced or absent in PNH. This led Dacie to suggest that PNH was an acquired clonal disorder arising in a haemopoietic stem cell. This important and perceptive idea was later confirmed by an elegant report in two patients with PNH who were also heterozygous for the enzyme glucose-6-phosphate dehydrogenase. In this study, it was shown that the patients’ red cells with a PNH phenotype contained only one isotype of glucose-6-phosphate dehydrogenase whereas their residual, non-PNH cells contained both.

Abbreviations: AA, aplastic anaemia; ER, endoplasmic reticulum; GPI, glycosyl phosphatidylinositol; GlcNAc, N-acetylglucosamine; FEA, phoshophoethanolamine; PI, phosphatidylinositol; PNH, paroxysmal nocturnal haemoglobinuria; VSG, variant surface glycoprotein
"Ham’s test was the diagnostic gold standard until its replacement by flow cytometry in recent years".

From the 1960s onwards, an increasing number of proteins were shown to be missing from the cell surface in PNH. These included molecules known to be involved in the regulation of complement at cell surfaces. It was hypothesised that their absence caused a complement mediated intravascular haemolysis. Decay accelerating factor (DAF/CD55), which has a role in the inactivation of complement at an early stage of the cascade, was thought to be important but individuals with the Inab red cell phenotype, who have an inherited DAF deficiency, had no clinical illness or in vitro red cell complement sensitivity. 12 CD59, which inhibits the formation of the membrane attack complex (the final step in the complement cascade) was the next candidate. Clinical evidence for the importance of this molecule came in 1992 when a 22 year old man was described who had a homozygous deficiency of CD59 on all his cells and suffered PNH-like symptoms, with haemolysis and cerebral thrombosis. 13 It is probable that the haemolytic and thrombotic features of PNH are mediated by complement sensitivity and that CD59 deficiency is an important cause of this.

The biochemical explanation for the absence of these proteins became clear when the GPI anchor was described as a novel mechanism of attachment of antigens to cells in 1980. 14 It was subsequently shown that all the proteins absent in PNH were GPI linked and that all GPI linked antigens are missing from PNH cells. In the past decade, the structure and biochemistry of the GPI anchor have been described and there is a consistent biosynthetic abnormality in all patients with PNH described to date. 15 A gene whose cDNA is able to correct this defect in all transfected human cell lines has now been cloned. 16 It is situated on the short arm of the X chromosome and has been named PIG-A, a term derived from its ability to restore the GPI synthetic defect in class A murine cell lines. 17 Since the gene has been cloned, mutations have been found in all patient samples reported. 18 This sequence of historical milestones has taken PNH from its most obvious clinical manifestation right back to a single gene defect in a haematopoietic stem cell. The story is in one sense complete but there are still many intriguing questions to be answered.

Figure 1  An example of peripheral blood phenotyping in paroxysmal nocturnal haemoglobinuria (PNH). Plots from a flow cytometer showing the clear discrimination between populations of normal and PNH cells in a patient’s peripheral blood sample. (A,B) Plots showing the three types of red blood cells that can be detected in these patients. Type I cells are normal, type II are partially deficient in glycosyl phosphatidylinositol (GPI) linked proteins, and type III are wholly deficient. CD55 and CD59 are GPI linked antigens found on red blood cells. (C) Plot showing granulocytes that have been dual stained with two GPI linked surface antigens (CD16 and CD66) in a patient with PNH. This clearly delineates the normal and the PNH cells. (D) Plot using the same double staining method but with antigens relevant to monocytes (CD64 and CD14): the same clear demarcation is shown. Courtesy of Dr S Richards, Haematological Malignancy Diagnostic Service, The General Infirmary, Leeds, UK.
nosis was 10 to 15 years in two large historical studies. The disease does shorten life. The median survival from diagnosis is 4 years and the overall frequency is probably similar to that of AA and only laboratory evidence of PNH are much less likely to suffer a thrombosis than those with active haemolysis and a large proportion of PNH cells in their blood. In this last group, thrombosis may occur in up to 50% and be the cause of death in one third. The intimate connection between AA and PNH is underlined by the clinical course of the illness in individual patients. Some degree of cytopenia is a consistent finding, even in haemolytic PNH. This may range from a mild reduction in one cell lineage to life threatening bone marrow failure. Even when blood counts are normal, bone marrow examination and progenitor culture assays reveal impaired haemopoiesis. Malignancy, such as myelodysplasia or acute myeloid leukaemia supervenes in around 5% of cases, but is probably a result of the process leading to AA rather than a specific risk related to the PNH clone itself, which is not considered preleukaemic.

Diagnosis
The demonstration of non-immune haemolysis with haemoglobinuria should lead to an investigation for PNH. Alternatively, its presence may be sought because of AA or a venous thrombosis at an unusual anatomical site. The diagnosis is definitively established by the demonstration of GPI linked protein deficiencies on red blood cell and neutrophil surfaces by multiparameter flow cytometry. The Ham test has been largely abandoned where flow cytometry is available, because it is relatively insensitive and labour intensive and only gives information on red blood cells. The solid phase gel techniques used for antibody detection in blood transfusion provide a rapid screen but again are limited to red blood cell antigens. The proportion of affected red blood cells often gives a falsely low assessment of the true clone size because of the effects of transfusion. The proportion of normal counterparts and because of the effect of transfusion. The Neutrophil series is not affected by these variables and therefore allows an accurate measurement of the clone size. It is possible to detect PNH clones that comprise < 1% of neutrophils using multiparameter flow cytometry. It is important to include a transmembrane antigen as a lineage marker (for example, CD15 for neutrophils) and at least two GPI linked antigens (for example, CD55 and CD59) to exclude the rare inherited deficiencies of single antigens such as the Inab phenotype (CD55 deficiency).

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Using flow cytometry, it is possible to demonstrate patterns of complete or partial GPI linked protein deficiency on the red blood cell series. Normal cells are designated type I, partially deficient type II, and completely deficient type III (fig 1). The clinical severity of the disease is directly related to the proportion of type III red blood cells. If these techniques are applied to other haemopoietic cell lineages, GPI deficiency can be documented on platelets, monocytes, and lymphocytes, confirming the stem cell nature of the disorder (fig 1).

Figure 2. The glycosyl phosphatidylinositol (GPI) anchor. This is a simplified diagram of the GPI structure. The C-terminus of the anchored protein links to an ethanolamine residue on the GPI anchor. The anchor itself consists of this ethanolamine moiety attached to a glycan core. The GPI structure attaches to the cell membrane via phosphatidylinositol. The glycan core consists of a molecule of GlcNAc linked to three mannose residues. The first step in GPI synthesis is the linkage of the GlcNAc to PI. It is this reaction that fails in paroxysmal nocturnal haemoglobinuria because the genetic lesion disrupts the production of a necessary enzyme complex.

CLINICAL ASPECTS OF PNH
Epidemiology
PNH is a rare disease. One of the largest epidemiological studies looked back at data from French centres from 1946 to 1995 and found only 220 reported cases. The annual incidence is about 0.4/million and the overall frequency is probably similar to that of aplastic anaemia (AA) with which it has a close association. It is probable that greater awareness and improved diagnostic methods will increase the number of diagnosed cases. The UK PNH registry in Leeds has been collecting new and existing cases since 1990 and currently has over 140 recorded patients (P Hillmen, personal observation, 2001).

Clinical features
Patients with PNH may have a long term chronic illness but the disease does shorten life. The median survival from diagnosis was 10 to 15 years in two large historical studies. Patients most commonly die of thrombosis or progressive cytopenias. Leukaemic transformation is uncommon (< 5%). Many patients will continue to have intermittent paroxysms of haemolysis but some eventually achieve a spontaneous remission. The identification of patients destined to remit is clearly an important requirement to prevent the use of toxic treatments in patients with a good prognosis.

Haemolysis is the cardinal feature. It is classically paroxysmal and most apparent in the first urine passed on waking—hence the name of the disease. In practice, patients often have a chronic haemolysis with exacerbations. This results in a variable transfusion requirement with iron deficiency often contributing to the anaemia. All patients should receive daily folic acid, because a low degree of haemolysis is usual between paroxysms. Heavily transfused patients can, paradoxically, become iron overloaded and this should be monitored to avoid compounding the problem with iron supplements.

Thrombosis is the most feared complication. There is a well established predilection for the hepatic veins but a diversity of predominantly venous sites has been described. Patients with AA and only laboratory evidence of PNH are much less likely to suffer a thrombosis than those with active haemolysis and a large proportion of PNH cells in their blood. This last group, thrombosis may occur in up to 50% and be the cause of death in one third.

The intimate connection between AA and PNH is underlined by the clinical course of the illness in individual patients. Some degree of cytopenia is a consistent finding, even in haemolytic PNH. This may range from a mild reduction in one cell lineage to life threatening bone marrow failure. Even when blood counts are normal, bone marrow examination and progenitor culture assays reveal impaired haemopoiesis. Malignancy, such as myelodysplasia or acute myeloid leukaemia supervenes in around 5% of cases, but is probably a result of the process leading to AA rather than a specific risk related to the PNH clone itself, which is not considered preleukaemic.

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Treatment
This is another area much influenced by the interplay of PNH and AA. Those with AA or progressive pancytopenia may be candidates for intensive disease modifying treatment, including immunosuppression or bone marrow transplantation. These approaches are usually not appropriate for classic PNH without bone marrow failure. Interesting exceptions to this rule are disease occurring in patients with a syngeneic twin. In this circumstance, there is little risk from transplant, although it appears that at least moderate doses of conditioning chemotherapy before stem cell infusion are necessary to prevent re-expansion of the PNH clone.20

Most patients without pronounced cytopenias simply require supportive management. Blood transfusion is the mainstay for those with symptomatic anaemia. Folate supplementation is mandatory and iron status should be monitored because patients can become iron deficient through urinary loss or iron overloaded from blood. Because thrombosis is a leading cause of mortality in this group, all those with haemolytic disease or a large percentage of PNH cells in their blood (perhaps > 50% PNH neutrophils) should be considered for primary prophylaxis with warfarin if there are no contraindications.21

THE BIOCHEMICAL AND MOLECULAR BASIS OF PNH
A failure of GPI anchor synthesis is a constant and key feature in all cases of PNH. The genetic basis of this abnormality is now well described, as is the detail of the biochemical consequences.16–20 GPI deficiency causes a loss of many proteins from the cell surface. The resulting cell phenotype explains the clinical features and suggests a mechanism for expansion of the PNH clone. These assumptions, while reasonable, await further experimental proof and the correlation of all clinical sequelae with specific protein loss has yet to be achieved. Before speculating on this, it is worth describing what we know about the GPI anchor, the missing proteins, and the underlying genetic defect.

The GPI anchor
Most cell surface proteins are attached via a sequence of hydrophobic amino acids that spans the lipid portion of the bilayer. This transmembrane domain achieves a stable interaction, which is not easily disrupted without destroying the membrane itself. In the 1980s, another method of cellular attachment was described whereby proteins were linked to a GPI molecule, which was itself inserted into the cell membrane.21 The structure and the biosynthesis of this GPI anchor were determined from work in trypanosomes, whose major surface glycoprotein is attached by this mechanism.22 The backbone of the GPI structure is highly conserved between species.

There are essentially three parts to the GPI anchor (fig 2). The membrane attachment is achieved through the insertion of the lipid moiety of phosphatidylinositol (PI) into the outer leaflet of the membrane. There is then a glycan core consisting of a molecule of N-acetylgalactosamine (GlcNAc) linked to three mannose residues and then to an ethanolamine. The protein attachment site is to the phosphoethanolamine (PEA) molecule linked to the terminal mannose. The C-terminus of the relevant protein is linked to the amino group of the PEA molecule by an amide bond.23–25

"Glycosyl phosphatidylinositol deficiency causes a loss of many proteins from the cell surface"

Biosynthesis of the GPI moiety occurs in the rough endoplasmic reticulum (ER). The precise location at which each reaction takes place is still a matter of some doubt. Some of the steps take place on the cytoplasmic surface of the ER and some within the cisternal space.26 The first step in the process is the addition of a molecule of GlcNAc to a PI residue. This does take place on the cytoplasmic surface of the ER and, at some point, the developing molecule is flipped to the cisterna. A series of three mannosylations follows using dolichol-phosphate mannose as a donor. Ethanolamine is added to each of these sugars. Transamidation of the newly synthesised protein leads to its attachment to the PEA molecule on the terminal mannose and finally the protein–GPI complex is transported to the external surface of the cell.

The importance of GPI linkage
The high degree of interspecies conservation of the GPI structure and its wide distribution argues for an important biological role for this mechanism of protein attachment. Most available information comes from work on trypanosomes and animal studies and the relevance to humans is speculative.

Enzymes have been characterised that cleave GPI anchors, thus releasing the tethered proteins.23 These phospholipases are present in other mammals and trypanosomes and their existence suggests that some proteins may be anchored through GPI to allow their selective removal. An example of this is the deliberate cleavage of the major surface protein (variant surface glycoprotein; VSG) of the trypanosome and its replacement with an immunologically discrete variant VSG from its repertoire. This allows the organism to evade the immune response in an infected host.23

Proteins attached through GPI are less tightly bound than their transmembrane counterparts, which allows a degree of transfer from one species (or cell) to another, as has been described in the parasitic infection caused by Schistosoma mansoni. In this case, the parasite seems to acquire host CD55, which in turn helps it to avoid complement mediated immune attack.23

In addition to allowing “loss or gain” of proteins, the biochemical membrane associations and mobility are different for GPI anchors and transmembrane domains. It is possible that the anchor’s characteristics and localisation are integral to the normal function of the associated protein.

GPI linked proteins are not randomly distributed over the cell membrane. In polar cells they are frequently located at the apical pole. In all cells, they associate with each other in regions of the membrane that are rich in glycolipids (sphingolipids and cholesterol), in so called glycolipid rafts. The importance of these structures remains unclear.

The GPI abnormality in PNH
Affected cells in PNH synthesise little or no GPI anchor. This results from a failure in the first step in the synthetic process—the addition of GlcNAc to PI. This has been demonstrated using different techniques by several workers. Biochemical studies using labelled precursors show an almost complete lack of intermediates containing mannose or glucosamine, indicating that the block in the pathway is at the first step.22 Murine cell lines incapable of synthesising GPI structures have been studied. In these experiments it could be shown that lines in which the defect occurred at different points in the pathway complemented one another (restored the synthesis when fused together). Three lines, classes A, C, and H, were individually unable to add GlcNAc to PI but when fused together they complemented one another.27 This implied that there were at least three gene products controlling this step. When PNH cells from patients are fused with these lines, they always complement cells of classes C and H but never those of class A. Thus, it became clear that the defect in PNH was always the same as that found in class A cells and led to a failure in the first step in GPI synthesis.23 This led to the subsequent discovery and expression cloning of the gene involved, which was termed PIG-A (phosphatidylinositol glycan complementation class
A). It appears that the product of PIG-A, along with at least three other gene products (PIG-C, PIG-H, and hGP1), form the enzyme complex responsible for the transfer of GlcNAc to PI (R Watanabe et al. In: Proceedings of the International Symposium on Glycosyltransferases and cellular communication, 1997, Osaka, Japan, abstract 6).

**GPI linked proteins in PNH**

If no GPI molecule is produced then the unlinked proteins are degraded in the ER and are absent from the cell surface. Some can still be expressed in an alternative transmembrane bound form (for example, FcγRIII/CD16 or LFA-3/CD58), but whether they retain the same function is not clear. A further complicating factor in the analysis of surface phenotype in PNH is that some patients can synthesise small quantities of GPI anchor and there appears to be competition between proteins for this residue, leading to partial expression of certain molecules and complete absence of others. This is best observed in red blood cells that are divided into three types on this basis by flow cytometry. Type I are normal in their surface expression, type II show reduced but detectable amounts of GPI linked proteins, and type III are completely deficient. Patients with florid haemolysis usually have a large proportion of type III cells, whereas those with non-haemolytic PNH in association with overt aplasia may have either type II cells or small type III clones. Type II cells probably arise in patients with PIG-A mutations that allow a small amount of residual protein to be produced—for example, some missense point mutations.

A wide range of proteins use the GPI linkage mechanism for cell surface attachment. There is no obvious similarity between them, belonging as they do to different functional groups. They include complement defence proteins, enzymes, blood group antigens, adhesion molecules, cell receptors, and others of unknown function. If the proteins are normally expressed on haemopoietic cell lineages then they are absent in the cells of the PNH clone. Table 1 illustrates the diversity of proteins that have been described, although it is by no means exhaustive.

“There is clearly a link between the clinical features of PNH and certain specific proteins lost through GPI synthetic failure. Understanding this would require a detailed understanding of the pathogenesis of the disease, which is at present incomplete. It would also require a full knowledge of the function of the proteins that are absent, which is also beyond us at present. Despite these problems, there are some associations that are quite well characterised and serve to whet our appetite. Most prominent among these is the intravascular haemolysis from which PNH derived its name. This is caused by a lack of one or more complement defence proteins from the red blood cell surface, allowing inappropriate and unopposed activation of complement. The candidate molecules include CD59 and CD55. Evidence from clinical studies, antibody blocking experiments, and gene transfer protocols favours CD59 as the most important molecule in this process, but a contribution from others cannot be ruled out. The thrombotic tendency in PNH is less well understood but may also result from CD59 deficiency. In this scenario, complement activation on platelet surfaces leads to increased eicosanoid, which exposes phospholipid as a site for thrombin generation. The biological consequences of the loss of the myriad of other GPI linked proteins from cells is not clear. In many instances, because of an element of redundancy in most biological pathways, there may be no sequelae, and in others the abnormalities may be subtle. The cause of the most intriguing feature of PNH clones—their relative growth advantage in a damaged bone marrow—remains obscure. It is possible that altered surface protein expression affects cellular responses or localisation within the microenvironment and the consequent alteration in the biology of the PNH cells allows them to evade the ongoing marrow insult and prosper in comparison with their normal counterparts. Our group has reported an example of abnormal localisation of haemopoietic stem cells in PNH. Most precursors in the peripheral blood of patients with PNH had the normal (non-PNH) phenotype, despite most bone marrow based progenitors being “PNH” in these individuals. We went on to show that treatment of these patients with granulocyte colony stimulating factor released largely PNH stem cells into the blood—altering the steady state. Elucidation of the specific (presumably GPI linked) mechanisms by which this and other cellular interactions occur is the Holy Grail of PNH research. The answers may also have much to tell us about haemopoietic stem cell biology, the pathogenesis of bone marrow failure states, and the pathophysiology of autoimmunity.

**The PIG-A gene**

In 1993, Miyata et al described the expression cloning of the PIG-A gene by the correction of a GPI deficient murine cell line, which had a similar GPI biosynthetic defect to that observed in PNH. The PIG-A gene is somatically mutated in all cases of PNH, presumably because it is the only gene of the GPI biosynthetic pathway that is found on the X chromosome at Xp22.1. This means that a single mutation of the gene on the active X chromosome of a haemopoietic stem cell in women or the only X chromosome in men, will result in the PNH phenotype. The PIG-A gene consists of six exons spanning 17 kb of genomic DNA. It has an open reading frame of 1432 bp encoding a protein of 484 amino acids. There is a short 5′ non-coding region with the initiation of transcription in exon 2 and a relatively large 3′ non-coding region. The
PIG-A promoter sequences are characteristic of a housekeeping gene, which presumably reflects the widespread expression of GPI linked proteins in all cell types.

**PIG-A mutations in PNH**

Because the mutations are somatic in PNH, they are extremely varied, with very few reported more than once. Approximately two thirds are small insertions or deletions resulting in a frameshift and early termination of transcription. In this circumstance, no active PIG-A product is produced and the PNH cells are completely deficient in all GPI linked proteins. The remainder are point mutations and these may result in a complete or partial deficiency of GPI linked proteins. More than 100 PIG-A mutations have now been described and very few are repeated (fig 3).

In over half of affected patients, flow cytometric analysis of the red blood cells identifies two discrete populations of PNH cells (type III, with complete deficiency; and type II, with partial deficiency of GPI linked antigens). This indicates that there are at least two unrelated PNH clones in these patients. Several groups have now described patients with more than one PNH clone at a molecular level; in fact, as many as four individual clones occur at the same time, but one patient was studied before and many years after a bone marrow transplant, at the time of relapse, and the PIG-A mutations at relapse were different to those of the original disease. In many of the cases with more than one mutated clone, red blood cell flow cytometry only identifies a single PNH population. Thus, it appears that most patients have multiple PNH clones. This indicates that patients are permissive for the development and/or expansion of PNH clones and therefore suggests a factor extrinsic to the PNH clone(s) that favours their development.

Is a PIG-A deficient clone sufficient to result in PNH?

The PIG-A gene is essential for embryogenesis and therefore mice that have “knocked out” PIG-A genes are not viable. When PIG-A deficient embryonic stem cells are microinjected into murine blastocysts, chimaeric mice are occasionally produced but only have a small number of cells derived from the PIG-A negative embryonic stem cells. The deficient stem cells contribute to the haemopoietic compartment of the resulting chimaeric mice but their proportion in any individual mouse remains constant with time. In addition, when mice are produced with higher proportions of GPI deficient haemopoietic cells, by using the Cre-Lox P system and/or by transplantation experiments, the proportion of GPI deficient haemopoietic cells remains constant over time. These findings show that PNH cells do not have a growth advantage over their normal counterparts in mice without bone marrow failure.

“The cause of the most intriguing feature of paroxysmal nocturnal haemoglobinuria clones—their relative growth advantage in a damaged bone marrow—remains obscure”

Araten and his colleagues recently reported the presence of rare GPI deficient neutrophils in normal individuals. These cells have a frequency of 10–51/million cells and when collected by flow sorting were shown to contain mutations of the PIG-A gene. These findings show that such mutations exist frequently among normal individuals, but this alone is not sufficient for the development of PNH.

**SUMMARY: THE DEVELOPMENT OF PNH**

We may now put forward a general outline of the factors leading to PNH and of its interplay with aplasia. The dual pathogenesis model appears to withstand the rigours of both time
and experimental research. In this hypothesis, somatic mutations in PIG-A lead to PNH only if the affected cells are in a bone marrow under hypoplastic stress. The mutation and the abnormal bone marrow environment are both required. PNH cells possess only a relative growth advantage and will not prosper in a normal bone marrow. If this is correct, then certain observations would be expected. Hypoplasia should be found invariably. This clinical association has of course been long recognised. Some degree of single or multiple cytopenia can be found in up to 80% of cases and in others there is laboratory evidence of diminished progenitor growth potential. It appears then, that hypoplasia and PNH do indeed go hand in hand. For dual prosper in a normal bone marrow.

The mechanism by which aplasia imparts a relative growth advantage to PNH cells is more speculative. Although few doubt the immune component in aplasia, it is unclear whether this is a primary alteration in the stem cell pool against which an immune response develops or an autoreactive state, where otherwise normal stem cells are targeted. In either scenario, the PNH cells may prosper by evading the immune destruction. This could be mediated directly by loss of a GPI linked “recognition molecule” or occur through altered biology or localisation of the cells through GPI linked mechanisms. The answers to these questions will not only enlighten PNH research but may greatly enhance our understanding of aplasia and stem cell behaviour.

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Alterations of the MDV oncogenic regions in an MDV transformed lymphoblastoid cell line

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Aims: Lymphoblastoid cell lines derived from Marek’s disease virus (MDV) induced tumours have served as models of MDV latency and transformation. They are stable and can be cultured with no detectable MDV genomic alterations upon repeated passaging. An MDV transformed lymphoblastoid T cell line (T9 cell line) has been reported to contain a disrupted MDV BamHI-H fragment and a Rous associated virus insertional activation of the c-myc protooncogene. In an attempt to define the respective participation of c-myc and MDV in the transformed phenotype of T9 cells, an analysis of MDV oncogenic sequences (BamHI-H, BamHI-A, and EcoQ fragments) was performed in these cells.

Methods: Using two different passages of the T9 cell line (late and early passages), the organisation of the MDV oncogenic regions and their expression in these cells was analysed. In vivo assessment of the oncogenicity of the virus contained within these cells was assessed by injecting them into 1 day old chickens.

Results: In T9 cells maintained in culture for up to six months (late T9), the MDV ICP4 gene was disrupted, whereas the meq gene was actively transcribed. The alterations of the MDV genome in these cells correlated with the inability of the virus to induce the classic signs of Marek’s disease in 1 day old chickens. However, early T9 cells submitted to a limited number of passages induced classic MDV pathogenicity, as efficiently as the MDV control cell line (T5), and did not show gross structural changes in the oncogenic MDV sequences.

Conclusions: Although the expression pattern of the MDV oncogenes in early T9 cells was identical to the one reported for other MDV transformed cells, longterm culture of an MDV transformed cell line containing a RAV insertional activation of the c-myc protooncogene led to the disruption of the MDV BamHI-H and BamHI-A oncogenic regions. In the late T9 cells MEQ was the only detected MDV oncoprotein. These results suggest that in the late T9 cells the truncated MYB protein compensates for the loss of MDV oncoproteins and reinforce the possibility that MEQ and MYB cooperate in the maintenance of the transformed state and the tumorigenic potential of these cells.

M arek’s disease virus (MDV) is an avian herpesvirus that causes T cell lymphomas and mononuclear infiltration in peripheral nerves, leading to paralysis within four to six weeks in susceptible chickens.1 The MDV genome, co-linear with α herpesvirus genomes, is 180 kb long and consists of short and long unique regions (US and UL, respectively), flanked by terminal repeats (TR, and TR, respectively), and internal repeats (IR, and IR, respectively).2,3

The rapid onset of tumours suggests the presence of MDV encoded oncogene(s). However, little is known about the molecular mechanisms of MDV induced oncogenesis. Several experimental approaches have been taken to identify viral genes that may be involved in the process of oncogenesis. These approaches include the analysis of: (1) the regions of the MDV genome that undergo alteration during attenuation of oncogenic strains (serotype 1); (2) the viral gene products expressed in MDV induced tumours and tumours derived from lymphoblastoid cell lines; and (3) the viral gene products of the serotype 1 strain, which are absent in the non-oncogenic serotype 2 MDV and serotype 3 herpesviruses of turkey strains. At least 15 open reading frames (ORFs) mapping mainly within four BamHI fragments (BamHI-H, BamHI-L-Q2, and BamHI-A) spanning the IR, and the IR, are expressed in transformed lymphocytes.1

“Much evidence has accumulated to suggest that MEQ might play a role in oncogenesis: it is consistently expressed in most MDV infected cells, in transformed cell lines, and in CD4+ T cells from lymphomas.” Initially, it was noted that serial in vitro passages of oncogenic MDV strains in primary chicken embryo fibroblasts (CEF) led to a loss of tumorigenicity;15,16 This change was shown to be associated with the amplification (up to 100 copies) of a 132 bp repeat within the BamHI-H and BamHI-D fragments.16 In lymphoblastoid cell lines carrying attenuated MDV strains, this amplification results in the expression of truncated RNA species from the 1.8 kb family.17 Kawamura et al.11 have reported that oligonucleotides complementary to the predicted splice donor site in the 1.8 kb family inhibit proliferation of MDV cell lines.11 The transfection of CEFs with two cDNAs isolated from this family of transcripts resulted in reduced serum dependence and prolonged proliferation of these cells.12 Four alternatively spliced and unspliced RNAs spanning the 132 bp repeat have been identified in CEFs infected with an oncogenic MDV strain, and at least two ORFs have been characterised, although their function remains unclear.12,13 Another protein, the phosphoprotein pp38, is expressed from this region. Its gene is transcribed in the opposite orientation and overlaps the US and IR, junction. The

Abbreviations: CEF, chicken embryo fibroblast; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; IR, long internal repeat; IR, short internal repeat; LATs, latency associated transcripts; MDV, Marek’s disease virus; MHC, major histocompatibility complex; ORF, open reading frame; PBS, phosphate buffered saline; RAV-2, Rous associated virus 2; TAE, Tris acetate EDTA; TR, long terminal repeat; TR, short terminal repeat; US, short unique region; UL, long unique region.
BamHI-D fragment encodes a phosphoprotein, pp24, the transcripts of which overlap TR L and U L. Both of these proteins are expressed in lytically infected cells and at a lower degree in lymphoblastoid cell lines. The use of antisense strategies suggested that pp38 might play a role in the proliferation of the MDV transformed cell line, MSB-1.

The MDV ICP4 gene maps to the BamHI-A fragment, within the IR S flanking the U L region of the MDV. It encodes an immediate-early transactivator that plays a crucial role in the regulation of transcription during MDV replication. Oligonucleotides complementary to the translation start region of ICP4 and the expression of antisense RNA inhibited the growth of MSB1 cells, indicating that the ICP4 gene might be required for the maintenance of the transformed state. Latency associated transcripts (LATs) expressed in lymphoblastoid cell lines and lytically infected CEFs are complementary to ICP4. It has been suggested that the LATs could regulate the expression of ICP4 and thus contribute to transformation by preventing replication of the virus.

The best characterised candidate MDV oncogene is the EcoRI-Q protein (MEQ), which is expressed from the adjacent BamHI-I2 and BamHI-Q2 fragments in the IRL. Much evidence has accumulated to suggest that MEQ might play a role in oncogenesis: it is consistently expressed in most MDV infected

Figure 1  (A) Genomic structure of Marek’s disease virus (MDV). MDV consists of long and short unique sequences (UL and US, respectively), flanked by long and short internal repeats (IR and IR, respectively), and long and short inverted terminal repeats (TR and TR, respectively). (B) Locations of the BamHI-D, BamHI-H, BamHI-I2, BamHI-Q2, BamHI-L, and BamHI-A regions. A detailed restriction map is indicated [E, EcoRI; B, BamHI; P, PvuII]. (C) Locations of the pp24, pp38, meq, and ICP4 homologue genes. Corresponding gene regions are enlarged to show the open reading frame and the transcription sense (arrow); the localisations of the oligonucleotides (arrowhead) used for PCR and Southern blotting are indicated. A solid bar represents the resulting amplified product used as a probe.

Figure 2  Disruption of Marek’s disease virus (MDV) BamHI-H fragment in late T9 DNA. Southern blots of BamHI digested high molecular weight DNA (15 µg) were hybridised with the [32P] labelled MDV specific 5.4 kb BamHI-H probe. DNA samples from chicken embryo fibroblasts (CEF s) were used as negative control. Molecular weight markers (in kilobases) are from HindIII digested λ DNA.
cells, in transformed cell lines, and in CD4+ T cells from lymphomas.22–24 The expression of RNA complementary to meq can prevent the growth of the MSB1 cell line,24 and overexpression of meq in rat-2 fibroblasts by recombinant murine retrovirus led to cell transformation.25 The MEQ protein is a transactivating b-Zip protein that homodimerises or heterodimerises in vitro with c-Jun, c-Fos, and cAMP response element binding protein.26–28 It has antiapoptotic activities,29 and was proposed to promote cell cycle progression.30 However, MEQ alone is not capable of transforming primary CEFS and the injection of chickens with replication defective virus carrying meq yielded a low incidence of sarcomas (5%), which eventually metastasised into internal organs.30–32 Together, these observations indicated that none of the viral genes is sufficient on its own for the induction and/or the maintenance of transformation of lymphoid cells in Marek’s disease. The close relation seen in the regulation of their expression suggested a synergistic action between these different viral genes. The pp38 promoter contains an ICP4 responsive element and transfection of MSB-1 cells by an ICP4 expression plasmid correlates with overexpression of the pp24 and pp38 phosphoproteins.33 The MEQ homodimer can bind to at least two distinct motifs (MERE I and II). A MERE II element is present in the putative MDV origin of replication, which overlaps the bidirectional promoter of the pp38 gene and the 1.8 kb family gene.32

We have previously reported that, in addition to an altered MDV BamHI-H fragment, the MDV transformed T9 cells also contained a c-myb insertional activation. These results suggested that the maintenance of the transformed state was dependent upon the presence of a truncated c-MYB protein.34 To test this hypothesis, we analysed the putative oncogenes regions from the MDV in the T9 cell lines. Here, we report that, except for the meq gene, MDV regions encoding putative oncogenes are altered in T9 cells, therefore raising the possibility that maintenance of transformation in this cell line might require cooperation between the meq and truncated myb oncogenes.

**MATERIALS AND METHODS**

**Cells**

The T9 and T5 T cell lymphoma cell lines were isolated from two testicular tumours obtained after the injection of MDV (HPRS16 strain) into 1 day old white leghorn chickens. The injection of T9 and T5 cells into chickens induced Marek’s disease at the time the cell lines were established. T cell antigens have been characterised on the surface membrane of T9 cells35 and the T9 cell line has been used in cytotoxicity assays.36 The two other Marek’s disease cell lines were isolated in different laboratories: MDCC-MSB1 was derived from a splenic lymphoma37 and HPRS2 was derived from an ovarian lymphoma.38 The RP9 cell line was established from a Rous associated virus 2 (RAV-2) induced B transplacental lymphoma.39 These cell lines were grown in RPMI 1640 medium supplemented with 10% fetal calf serum. The BM2 myeloblastic cell line was obtained from a chicken infected with avian myeloblastosis virus.40 BM2 cells were grown in DMEM medium supplemented with 10% fetal calf serum, 10% tryptose phosphate broth, and 2% heat inactivated chicken serum. CEFS were prepared from 13 day old C/E brown leghorn embryos (gs+, chf+, V−) of the Edinburgh strain, and cultured as described elsewhere.41 All cultures were performed at 37°C in a 5% CO2 atmosphere.

**In vivo infectivity assays**

The GB1 strain of inbred chickens bearing the major histocompatibility complex (MHC) haplotype B13 was used in these studies.42 These chickens were free from specific pathogens, lacked maternal antibodies directed against MDV, and were highly susceptible to Marek’s disease. At hatching, three groups of chickens were injected intraperitoneally with 10^6 T5 or T9 cells. In each group, eight to nine chicks were injected and six were used as internal controls to assess the horizontal transmission of MDV. Birds were euthanised and necropsied upon signs of morbidity. Organs with Marek’s disease lesions were harvested, processed for histological examination, and frozen for subsequent DNA isolation. Histological examinations of the nerves were systematically performed.

**Southern blot analysis**

High molecular weight DNA was purified from primary cells, cell lines, and solid tumours as described previously.43 After restriction endonuclease (GIBCO-BRL, Cergy Pontoise, France) digestion and electrophoresis in horizontal 0.8% TAE agarose gels, DNA fragments were transferred on to Nytran membranes (Schleicher and Schuell, Dassel, Germany). Hybridisation to [32P] labelled probes was performed as described elsewhere.44 All probes were prepared from purified DNA inserts.

**RNA purification and analysis**

Total RNA was prepared from cultured cells, using the guanidinium isothiocyanate method and northern blotting analysis was performed under the conditions described previously.45

**Polymerase chain reaction**

DNA from whole cellular lysates, prepared as described,46 and purified high molecular weight DNA were subjected to polymerase chain reaction (PCR) amplification using Taq DNA polymerase (Appligene, Paris, France) in a Perkin-Elmer Cetus
The primers for the pp38 gene used for amplification were as follows (5′ to 3′): pp38SD, GCTGCAGCTGTCCATTTTCC; pp38SI, GCCATCCTTGTCTTTCTGCC; and pp38I, CTGCTTCGAATTCCATCACC (nt 1529 to 1549, 2564 to 2584, and 2500 to 2519, respectively). The primer for the pp24 gene was as follows: pp24SI, ACCCCGTAACCAGCATGATG (nt 1411 to 1431).

The primers for the ICP4 gene were as follows: ICP4SI, GCCATGGGATGTGTTGAATCT; ICP4SD, CAACGCCAATATGGACGATGA; and ICP4B, TGAGACTTCACCGTCAAATG (nt 1395 to 1414). The PCR conditions were as described previously. The thermocycler for PCR was as described previously. The primers for the pp38 gene used for amplification were as follows (5′ to 3′): pp38SD, GCTGCAGCTGTCCATTTTCC; pp38SI, GCCATCCTTGTCTTTCTGCC; and pp38I, CTGCTTCGAATTCCATCACC (nt 1529 to 1549, 2564 to 2584, and 2500 to 2519, respectively). The primer for the pp24 gene was as follows: pp24SI, ACCCCGTAACCAGCATGATG (nt 1411 to 1431). The primers for the ICP4 gene were as follows: ICP4SI, GCCATGGGATGTGTTGAATCT; ICP4SD, CAACGCCAATATGGACGATGA; and ICP4B, TGAGACTTCACCGTCAAATG (nt 1395 to 1414). The PCR conditions were as described previously.
The primers for the meq gene were as follows: MEcoQSD, ACATTGCTCCGGTTCCCAAC; MEcoQSI, ATAGACGATGTGCTGGCTGAG; and MEcoQ, TATGGGGAGTAGAGATGGGA (nt 843 to 862, 1392 to 1411, and 933 to 952, respectively). PCR products were separated on TAE agarose gels and analysed by Southern blot hybridisation with labelled internal oligonucleotide probes (pp38I, ICP4B, and MECoQ), as described previously.

Nucleic acid probes
DNA fragments were GeneClean (Bio101) purified, and labelled with $[^{32}\text{P}]$ dCTP (ICN, Orsay, France) by nick translation (Invitrogen, Cergy Pontoise, France). The MDV specific probe corresponded to the BamHI-H fragment localised at the junction between the U and IR. The meq and ICP4 DNA probes were obtained from PCR amplified fragments using DNA isolated from the MSB1 cell line and specific oligonucleotides (fig 1). The PCR products were sequenced and cloned in the pMOSBlue plasmid vector (Amersham, Orsay, France). Avian glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as an internal standard to assess the amount of RNA present in each sample.

Immunoblot assays
Cells were lysed in RIPA buffer (50 mM Tris/HCl, pH 8.0, 150mM sodium chloride, 0.1% sodium dodecyl sulfate, 0.5% sodium deoxycholate, 1.0% NP40, and 5mM EDTA); 5 µg/ml of leupeptin, antipain, chymostatin, pepstatin; 0.4 TIU of aprotinin/ml (trypsin inhibiting units); and 2mM of aminoethyl-benzenesulfonyl fluoride (Sigma, Saint Quentin Fallavier, France). Lysates were clarified by centrifugation at 10 000 $\times$ g for 15 minutes at 4°C and supernatants were stored frozen at −70°C until analysis. Samples of protein extracts solubilised in Laemmli buffer were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis and electroblotted on to a PVDF membrane (Immobilon P; Millipore, Strasbourg, France). Filters were incubated overnight at 4°C in phosphate buffered saline (PBS) containing 4% non-fat dried milk and 0.05% Tween 20, before probing with the appropriate antiserum for one hour at 37°C. Filters were rinsed three times in PBS containing 0.05% Tween 20 for 10 minutes and then incubated with horseradish peroxidase conjugated goat anti-rabbit antibodies. The proteins were detected by the enhanced chemiluminescence method (ECL), under the conditions recommended by the manufacturer (Amersham).

RESULTS
Structure of the BamHI-H and BamHI-D fragments in the MDV T cell lines
The T9 cells were serially maintained in culture and passaged for up to six months. Both low number (early T9) and high number passage stocks (late T9) were used in our study.

When PCR was performed with the pp38SI and pp38SD primer set a 1.0 kb DNA fragment was amplified from MSB1, T5, and early T9 DNA (not shown). No amplification was observed with late T9 and CEF DNA. The BamHI-H and BamHI-D share the same sequence within TRL and IRL, amplification of the pp24 gene fragment could be performed with pp38SI and the pp24SI primer specific for the 3′ proximal coding region of the pp24 gene (fig 1). A 0.8 kb fragment was amplified from the DNA of all MDV cell lines (not shown). No amplification was observed for CEF DNA.
Southern blot hybridisation performed with the BamHI fragment H as probe detected a 5.4 kb and a 12 kb BamHI fragment in early T9, T5, HPRS2, and MSB1 DNA, corresponding to the BamHI-H and BamHI-D fragments, respectively (fig 2), but not in the CEFs or in RP9 RAV transformed cells. The 5.4 kb BamHI-H fragment was not detected in late T9 cells, although the 12 kb fragment was still present, a result in agreement with amplification of the pp24 gene (fig 2, right hand panel). These results indicated that the BamHI-D fragment containing the pp24 gene was present in both early and late T9 cell DNA, and that the BamHI-H fragment is disrupted in the late T9 cells.

Structure and expression of the region containing the ICP4 gene and LATs in the MDV cell lines

The IRS contains the 5′ part of the BamHI fragment A, which encodes the immediate–early ICP4 gene. A specific fragment of the ICP4 gene obtained by PCR was used as a probe (fig 1).

Northern blotting using the ICP4 PCR product as a probe revealed that transcripts of 0.72 and 10.0 kb were expressed in MSB1, HPRS2, T5, and early T9 cell lines (fig 3). As previously reported, these RNA species resulted from antisense transcription of the ICP4 locus. No ICP4 RNA species were detected from either late T9 cells or from cells that are not transformed by MDV (CEF, BM2, and RP9).

Hybridisation of EcoRI digested high molecular weight DNA with the ICP4 DNA probe detected a 5.8 kb DNA fragment in T5, MSB1, and early T9 cell lines (fig 4). No fragments were detected by this probe in late T9 cell DNA. In early T9 cells, this probe revealed an additional, but less intense band corresponding to a 6.8 kb DNA fragment. As expected from the previously published nucleotide sequence, the same probe detected a 2.3 kb fragment in the PvuII digested DNA from the T5, early T9, and MSB1 cell lines (fig 4, right hand panel).

Table 1  Induction of Marek’s disease following injections of T5 and T9 cell lines

<table>
<thead>
<tr>
<th>Cellular strains</th>
<th>Age at necropsy (weeks)</th>
<th>Chickens with macroscopic lesions/injected ones (%)</th>
<th>Chickens with macroscopic lesions/contact ones (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T5</td>
<td>11</td>
<td>4/8 (50)</td>
<td>4/6 (66)</td>
</tr>
<tr>
<td>Early T9</td>
<td>6–8</td>
<td>8/9 (88)</td>
<td>5/6 (83)</td>
</tr>
<tr>
<td>Late T9</td>
<td>6–13</td>
<td>0/9 (0)</td>
<td>0/6 (0)</td>
</tr>
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</table>
These results indicated that the 5′ BamHI-A fragment containing the ICP4 gene is also disrupted in late T9 cells. As revealed by the detection of an additional 6.8 kb fragment, the 5′ BamHI-A fragment is rearranged in early T9 cells. From the size of this fragment and the origin of the probe, the rearrangement in early T9 cells maps within the promoter of the ICP4 gene between the EcoRI and PvuII restriction sites (positions 11193 and 7750, respectively; numbering according to the previously published nucleotide sequence).

Expression of the meq gene in the MDV cell lines

The Eco-Q fragment of the MDV genome encompasses the 3′ end of the BamHI-I2 and BamHI-Q2 fragments. It contains an ORF referred to as meq.

In MDV transformed cells, Southern blotting performed with the meq probe detected a 5.1 kb and a 1.3 kb EcoRI fragment (fig 5, right hand panel), respectively. The sizes of these fragments were in agreement with the previously published nucleotide sequence.

Wilson et al have reported that the reduced expression of MDV gC in attenuated strains did not result from structural alterations within or near that gene, and suggested that gene(s) encoding MDV regulatory protein(s) that interact with the MDV gC promoter might be altered during the attenuation process. The Eco-Q region expression pattern was established following northern blot hybridisation with the meq DNA probe. At least five transcripts with sizes ranging from 1.4, 1.8, 2.1, and 2.5 to 3.1 kb were detected in all the MDV cell lines (fig 6). Only minor differences in the degree of expression of these different species could be seen among RNA from these cell lines. This result was consistent with previous work reporting the expression of both spliced and unspliced sense transcripts in addition to antisense transcripts from the BamHI-I2/Q2 region encoding MEQ.

Immunoblotting performed with a polyclonal MEQ antibody revealed proteins of 31, 54, and 57 kDa in all cell extracts (fig 7). An additional 74–80 kDa doublet was specifically detected in MSB1, T5, and the early and late T9 cell extracts. The apparent molecular weight of these MEQ proteins was higher than those described previously. Liu and colleagues reported that the MEQ protein migrated with an apparent molecular weight ranging from 50 to 75 kDa, depending upon the origin of the cell lines analysed. These discrepancies may result from post-translational modifications, such as phosphorylation. Alternatively spliced MEQ RNA species also contribute to the variation in apparent sizes of the MEQ proteins.

Pathogenicity of the MDV cell lines in vivo

Marek's disease can be experimentally induced by the injection of MDV lymphoblastoid cell lines into susceptible birds. To determine whether early and late T9 cells were still able to induce Marek's disease, 10^7 cells were injected intraperitoneally into 1 day old chickens. The T5 cell line was used as a positive control.

Chickens injected with either T5 or early T9 cells developed classic signs of Marek's disease (paralysis; table 1). Furthermore, contact exposed chickens developed signs of Marek's disease, therefore confirming the horizontal transmission of infectious virus. Necropsy frequently revealed macroscopic tumours and bursal and thymic atrophy in either inoculated or contact exposed chickens. Lymphoid tumours were localised mainly to the sexual organs, but occasionally arose in liver, kidney, heart, and proventriculi (table 2). The sexual localisation appeared to be a characteristic of the oncogenic HPRS16.
Molecular analysis of the tumours (or normal tissues) isolated from T5, early T9, and late T9 cell line inoculated and contact chickens. (A) Detection of Marek's disease virus (MDV) sequences in DNA from induced tumours. PCR amplification was carried out on DNA isolated from different tissues of chickens injected with T5 (964, 955, and 959), early T9 (479 and 480), and late T9 (971 and 980). The control without DNA template (lane C) was performed under identical conditions. Additional controls included DNA isolated from the early T9, late T9, and T5 cell lines. PCR products were run on a 1.5% agarose gel and visualised with an ultraviolet transilluminator. The positions of selected bands from a 1 kb ladder marker (BRL) are indicated on the left. (B) PCR products run on the gel were transferred on to a nitrocellulose membrane and then hybridised with $^{32}$P labelled oligonucleotides pp38I. (C) PCR amplification of the 5' RAV–c-myb junction. The same DNA templates as in (A) were subjected to 30 rounds of amplification using primer 1 (corresponding to c-myb exon 3 sequences) and primer 2 (corresponding to RAV gag sequences). PCR products were run on a 1.5% agarose gel and visualised by means of an ultraviolet transilluminator. The position of the PCR amplified 850 bp 5' junction fragment is indicated. Control experiments were performed under identical conditions without DNA (lane C).
obtained after serial passage of virulent MDV strains in reminiscence of the situation encountered in attenuated strains late T9 cell line to induce Marek's disease in chickens. confirmed that these alterations abrogated the capacity of the were rearranged in the late T9 cells. In vivo experiments regions were identical to those from the MSB1 prototype cell intact in the early T9 cells and expression patterns from these analysis revealed that all these fragments were apparently previously reported activation of c-myb in T9 cells originate from an expansion of the injected tumour cells or amplified product was confirmed by probing with the pp38I from late T9 cell injected chickens (fig 8A). The origin of the amplified product was confirmed by probing with the pp38I labelled oligonucleotide (fig 8B).

To establish whether tumours arising in T9 infected birds originate from an expansion of the injected tumour cells or from MDV induced transformation, we took advantage of the previously reported activation of c-myb in T9 cells and performed MHC typing. Using primers specific for the 5' myb-RAV junction in T9 cells, no products were amplified from DNA derived from either early T9 induced tumours or tissues from late T9 inoculated chickens (fig 8C). Positive and negative controls were performed with late and early T9 and T5 cells, respectively. PCR-single stranded conformational polymorphism experiments performed with different sets of primers specific for the YF gene from the MHC class I established that tumoral cells exhibited the same pattern as that of the recipient chicken. This pattern was different from that of T5, early T9, and late T9 cells (data not shown). These results indicated that tumour formation in injected animals did not result from the proliferation of early T9 or T5 cells, but from the transformation of chicken recipient lymphoid cells by MDV.

DISCUSSION

We have previously reported that the MDV transformed T9 lymphoblastoid cell line contained a RAV insertional activation of c-myb and a rearranged MDV genome lacking the BamHI-H fragment. To establish the relative participation of the MDV and myb sequences in the transformed phenotype of the T9 cells, we analysed them for the presence of MDV oncogenic sequences. We have focussed our analysis on the following potential oncogenic regions: the BamHI-H, BamHI-A, and Eco-Q fragments, which encode the pp38 and the 1.8 kb RNA family; the ICP4 homologue transcriptional factor; and meq. Genomic analysis revealed that all these fragments were apparently intact in the early T9 cells and expression patterns from these regions were identical to those from the MSB1 prototype cell line and T5 cells, which express infectious transforming MDV. Conversely, the BamHI-H fragment and the ICP4 coding region were rearranged in the late T9 cells. In vivo experiments confirmed that these alterations abrogated the capacity of the late T9 cell line to induce Marek’s disease in chickens.

The loss of the BamHI-H fragment in late T9 cells is reminiscent of the situation encountered in attenuated strains obtained after serial passage of virulent MDV strains in primary CEF. Because no alteration of this region occurred upon repeated passages of the early T9 cells (not shown), the rearrangement of this region in late T9 cells must have involved another mechanism. Moreover, an additional, less intense fragment was revealed with the ICP4 probe in the early T9 cells that could result from rearrangement in progress in a subpopulation of cells. Hayashi and colleagues have previously described an amplification of a 178 bp repeat sequence within the 1.6 kb HindII subfragment of BamHI-A in viral DNA isolated from both pathogenic and non-pathogenic strains, and in established lymphoblastic cells. In this last case, the 178 bp expansions mapped about 1 kb downstream of the stop codon of the ICP4 gene, giving rise to heterogeneity of IR, and TR. In contrast, alteration in early T9 cells seems to take place in the ICP4 promoter region without revealing heterogeneity accounted for expansions. The expression of ICP4 is down regulated by the LATs during latency and the balance between the sense (ICP4) and antisense transcripts may serve as a switch to turn off MDV replication during latency. The lack of the ICP4 and LAT coding regions in the late T9 cell line may totally abrogate reactivation of the virus. Because an ori sequence is found in the bidirectional promoter of pp38 and the 1.8 kb family RNA gene (BamHI-H), it is likely that in the late T9 cells MDV is unable to replicate. Indeed, immunodiffusion performed with blood samples from chickens inoculated with late T9 cells did not reveal antibody against MDV (data not shown), confirming the fact that late T9 cells did not produce infectious viruses and were no longer capable of inducing Marek’s disease.

“Because MEQ was the only MDV oncogene fully expressed in the late T9 cells, our results raise questions regarding the factors involved in the maintenance of the transformed phenotype in these cells.”

Deletions and expansions of tandem repeats were reported to occur concurrently with attenuation of oncogenic MDV strains upon extensive passages in primary CEFs. Usually, MDV lymphoblastoid cells are immortalised cell lines, which are latently infected with MDV, and capable of transferring MDV to CEFs or duck embryo fibroblasts in vitro. Despite many years in culture, these cell lines are still able to induce Marek’s disease in susceptible birds. Repeatedly passaged T9 cells might have selected cells containing deletion of the BamHI-H and the ICP4 encoding region in their MDV genome and other gross structural alterations that were not detected in our study. The relative instability of the MDV genome in these cells could result from insertions, deletions, or other events mediated by RAV-1 and homologous recombination between long terminal repeat sequences because it has been reported that the MDV genome often undergoes structural changes, including insertions of retroviral elements. Identification and sequencing of junction sites at the border of these deletions could help in elucidating the mechanism involved in the disruption of the MDV sequences in late T9 cells.

Because MEQ was the only MDV oncogene fully expressed in the late T9 cells, our results raise questions regarding the factors involved in the maintenance of the transformed phenotype in these cells. We have previously shown that T9 cells expressed high amounts of a truncated MYB protein as a result of RAV-1 integration within the c-myb allele. A similar truncation conferred a transforming potential on myb, when assessed in primary chicken fibroblasts, and related chimaeric mRNA was isolated from B lymphomas and fibrosarcomas induced by avian leukosis virus. To our knowledge, this is the only report of a RAV insertional activation of c-myb in a T lymphoma. Liu et al reported that MEQ can transform established Rat-2 fibroblasts but MEQ alone is not capable of transforming primary cells. Furthermore, recombinant retroviruses carrying meq yield a low incidence of sarcomas, indicating that meq is a weak oncogene, and that meq may require additional cooperating oncogenes to display its full transforming mechanism. At least four MDV genes (pp38, ICP4, meq, and the 1.8 kb RNA family) have been shown to be important for the maintenance of the transformed phenotype.
in MSB1 cells. Because meq expression is detected in late T9 cells, we speculate that the activation of c-myc might compensate for the lack of MDV genes and act synergistically with meq to maintain the transformed phenotype in late T9 cells.

To determine whether the expression of both the MEQ and truncated MYB proteins is required for the maintenance of the transformed phenotype, we have engineered T9 cells that express antisense mRNA using the pMDL7 inducible vector (pMD221 for meq antisense and pMDMyb, our construct). In contrast to Morgan's laboratory, we failed to establish antisense expressing cell lines, but we were able to establish control cell lines (expressing sense MEQ and MYB). These results suggested that the inhibition of meq or myb expression might interfere with T9 cell proliferation; an observation in favour of a role for both the MEQ and MYB oncoproteins in the maintenance of the transformed phenotype. Another antisense strategy should be explored to determine the subtle contribution of activated myb and/or meq in this phenotype. Nonetheless, our results indicate that T9 cells constitute a cellular model to identify the meq target genes in the context of cells carrying a defective MDV genome.

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