INTRODUCTION

Membrane-bound proteins can be post-translationally modified by fatty acids (myristic and palmitic), isoprenoids (farnesyl and geranylgeranyl) or glycosylphosphatidylinositol (GPI) [1]. Modification of cell surface proteins by GPI serves as a ubiquitous anchoring mechanism that is found in parasites, yeast and mammals [2]. The glycan core of GPI-anchored proteins consists of the GPI anchor generally involves alkylacylglycerols or diacylglycerols, but in yeast and the slime mould Dictyostelium discoideum, ceramide-based GPls are also known [2]. Glucosaminyl(acyl)phosphatidylinositol [GlcN(acyl)PI], the third intermediate in the mammalian glycosylphosphatidylinositol (GPI) anchor pathway, is undetectable in most cells. This intermediate was previously shown to accumulate, however, in murine lymphoma mutant E and in yeast mutant dpm1, both of which lack dolicholphosphomannose synthase activity. Here we report that a mammalian HeLa S3 subline, denoted D, produces large amounts of GlcN(acyl)PI. The level of GlcN(acyl)PI in this subline is twice that in the murine lymphoma mutant E and 4 times that in the parental S3 line. This HeLa D subline differs from the previously reported mutants that accumulate GlcN(acyl)PI because no defects in the synthesis or utilization of dolicholphosphomannose were found. Kinetic analysis indicated that in this HeLa subline there is an increased rate of synthesis of GlcN(acyl)PI, whereas the rate of metabolism for this GPI is comparable to that in wild-type cells. Furthermore, HeLa D cells accumulate GlcN(acyl)PI without a block in the synthesis of the downstream mannosylated GPI anchor precursors and GPI-anchored proteins. These findings might be relevant for understanding the regulation of the GPI pathway.

GPI-anchored proteins. In contrast, this HeLa S3 subline synthesizes decreased amounts of downstream intermediates or the GPI intermediate preceding the site of the lesion and the synthesis and metabolism of H6, H7 and H8 seem to be in excess of what is needed to anchor GPI-anchored proteins [9]. Furthermore, kinetic analysis of the synthesis and metabolism of H6, H7 and H8 argues for roles in addition to that of precursors of protein anchors [10]. This hypothesis was further supported by the presence of these GPls outside the endoplasmic reticulum (ER), including in the plasma membrane [10–12].

GlcN(acyl)PI is present at very low levels in wild-type cells. It can be detected in the murine lymphoma mutant E [7] and the yeast mutant dpm1 [13] because both mutants are deficient in dolicholphosphomannose (Dol-P-Man) synthase, the enzyme responsible for the synthesis of Dol-P-Man. Dol-P-Man is the donor of the mannosyl residues in GPls as well as of four mannosyl residues in the lipid-linked oligosaccharide precursor of asparagine-linked glycan chains, Glc₃Man₃GlcNAc₂, -PP-Dol. Lec35, a Chinese hamster ovary (CHO) mutant with normal levels of Dol-P-Man but with a mutation affecting its utilization, was also suggested to produce high amounts of GlcN(acyl)PI and to give inefficient GPI mannosylation [14].

We recently reported the isolation and chemical characterization of GlcN(acyl)PI [6] in an S3 HeLa subline. In this study we compare the levels of GlcN(acyl)PI in four HeLa cell lines: CCL-2, S3 (a clonal derivative from the parental CCL-2), D (a S3-derived cell line) and D3 (a clone from D). Mutants with lesions in the GPI anchor pathway characteristically accumulate the GPI intermediate preceding the site of the lesion and synthesize decreased amounts of downstream intermediates or GPI-anchored proteins. In contrast, this HeLa S3 subline

Abbreviations used: CHO, Chinese hamster ovary; DAF, decay accelerating factor; DMEM, Dulbecco’s modified Eagle’s medium; Dol-P-Man, dolicholphosphomannose; EthN-P, ethanolamine phosphate; ER, endoplasmic reticulum; GlcN(acyl)PI, glucosaminyl(acyl)phosphatidylinositol; GPI, glycosylphosphatidylinositol; PI, phosphatidylinositol; PI-PLC, PI-phospholipase C; PIP, phosphatidylinositol phosphate; lysoPI, 2-lysophosphatidylinositol.

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accumulates GlcN(acyl)PI without significantly affecting the levels of downstream intermediates or GPI-anchored proteins.

**EXPERIMENTAL**

**Materials**

Dulbecco’s modified Eagle’s medium (DMEM) as well as inositol-free and glucose-free DMEM were obtained from Gibco. Heat-inactivated newborn calf serum was from Sigma. PI-phospholipase C (PI-PLC) was purified from the medium of a Bacillus subtilis strain that overexpresses PI-PLC of *B. thuringiensis* [15]. myo-[2-³H]inositol (20 Ci/mmol) was from American Radiolabeled Chemicals and [²-³H]Man (20–30 Ci/mmol) from ICN.

**Cells and culture conditions**

Two HeLa cell lines, CCL-2 and S3, were obtained from the American Type Culture Collection. S3 is a clone of CCL-2 that readily grows in suspension culture. Two additional lines, D and D3, were isolated in our laboratory. HeLa D was derived from S3 by continuous passage of HeLa S3 cells with a rubber ‘policeman’ or a bent Pasteur pipette to wash off the cells from the flask walls, whereas CCL-2 and S3 cells were routinely passed with trypsin. HeLa D3 is a clone of D obtained by limiting passage at 37 °C.

**Labelling and extraction of glycolipids**

Cells (10⁶) were plated in 75 cm² flasks and incubated with 10 μCi of [³H]inositol for 3 days in 10 ml of inositol-free DMEM supplemented as described above. For labelling with [³H]Man, 2 x 10⁶ cells were preincubated for 1 h in 2 ml of glucose-free DMEM supplemented with 100 μg/ml penicillin, 100 μg/ml streptomycin, and 2 mM glutamine in an atmosphere of 5 % CO₂ in air at 37 °C. Karyotype analysis of D cells showed the same modal chromosome count (65–67) and the same copy numbers of HeLa markers (one of M1, one of M2, two of M3, one of M4) reported for HeLa S3.

**Analysis of lipid-linked oligosaccharides by size-exclusion chromatography**

Cells (2 x 10⁶) were incubated with 1 mCi of [³H]Man in 1 ml of glucose-free DMEM supplemented with 10 % (v/v) newborn calf serum/100 μg/ml glucose for 20 min at 37 °C. Cells were then washed with PBS and resuspended in PBS/newborn calf serum (1:1, v/v), and chloroform/methanol (3:2, v/v) was added to give chloroform/methanol/water (3:2:1, v/v). The organic phase was back-extracted with aqueous phase pre-equilibrated with organic phase, dried and incubated in 500 μl of n-propanol and 1 ml of 10 mM HCl for 20 min at 100 °C. Hydrolysed samples were partitioned between chloroform and water, and 3 x 10⁶ c.p.m. of the water phase was subjected to chromatography on a P4 column. The relative elution constant (Kᵣ) of an oligosaccharide peak was calculated from the formula $Kᵣ = (Vᵣ − Vₑ)/(Vₑ − Vᵢ)$, where $Vᵣ$ is the elution volume (fraction number) of the oligosaccharide of interest, $Vₑ$ is the exclusion volume (determined with albumin) and $Vᵢ$ is the inclusion volume (determined with [³H]Man).

**SDS/PAGE**

GPI-anchored proteins were detected after extraction of the cells with Triton X-114 and treatment of the extracts with PI-PLC [18]. Cells labelled with 100 μCi of [³H]inositol for 3 days were extracted by incubation with 700 μl of 20 mM phosphate buffer, pH 6.9, containing 5 mM EDTA, 1 mM EGTA, 150 mM NaCl, 2 % (w/v) Triton X-114, 1 μg/ml leupeptin and 1 μg/ml pepstatin for 30 min on ice. The extract was centrifuged for 10 min at 15000 g. The pellet was re-extracted with 300 μl of the same buffer and the supernatants were combined. After phase separation at 37 °C [19] and centrifugation, the aqueous phase was removed and the detergent phase was washed with 300 μl of fresh aqueous phase. The pooled aqueous phase was then back-extracted with fresh condensed detergent phase, divided into two, diluted in 20 mM phosphate buffer and incubated overnight with PI-PLC (10 μg/ml) at 4 °C. At the end of the incubation period, the samples were warmed at 37 °C for phase separation. The detergent and the aqueous phases were washed by a second cycle of phase partitioning before mixing with 0.5 vol. of tri-chloroacetic acid (1 g/ml) containing 1 mg/ml sodium deoxycholate. After incubation overnight at 4 °C, the precipitates were collected by centrifugation at 14000 g for 15 min and washed twice with cold acetone. The pellets were resuspended in loading buffer and resolved by SDS/10% PAGE [20]. The gel was soaked in Amplify (Amersham) and subjected to fluorography.
Two-site immunoradiometric assay

Cells (2 × 10^6) were plated in 100 mm dishes the day before the experiment. On the next day the medium was removed and the resulting subconfluent cells (4 × 10^6) were washed once with DMEM and then incubated in 3 ml of DMEM containing 10 μg/ml PI-PLC for 30 min at 37 °C. A 1 ml portion was removed and centrifuged for 10 min at 15000 g, and the immunoradiometric assay was performed on the supernatant as previously described [21]. Briefly, microtitre plate wells were coated with anti-DAF monoclonal IA10 (capturing antibody), and blocked with 1 % (w/v) BSA in PBS, after which 25 μl of sample was added. After incubation for 2 h at 20 °C, 125I-labelled anti-DAF monoclonal IIH6 (revealing antibody, directed against a different DAF epitope) was added. Wells were then washed and the radioactivity remaining in the cut-out wells was measured in a γ-counter.

Modelling of radiolabelled inositol incorporation and decay

A simple kinetic model for inositol incorporation into the GPI biosynthetic pathway is shown in Scheme 1. In this simplified model, inositol (A) is incorporated into PI (B) with a rate constant k1. PI is modified further into GlcN(acyl)PI (C) with a rate constant k2, and GlcN(acyl)PI is converted into a mannosylated GPI (D) with a rate constant k3. The rate constants for degradation of each species are kα, kβ and kγ, respectively. The amounts of each species at the start are [A]0, [B]0 and [C]0 respectively.

\[
\begin{align*}
[A] & = [A]_0 e^{-\alpha t} \\
[B] & = k_1 [A]_0 (e^{-\alpha t} - e^{-\beta t})/(\beta - \alpha) + [B]_0 e^{-\beta t} \\
[C] & = k_2 [B]_0 (e^{-\beta t} - e^{-\gamma t})/(\gamma - \beta) + [C]_0 e^{-\gamma t} \\
[D] & = k_3 [C]_0 e^{-\gamma t}
\end{align*}
\]

Experimental data were fitted to eqns. (1)-(3) by nonlinear regression analysis with Fig.P (version 6.0), and parameter values that fitted both the pulse and chase initial conditions were estimated.

RESULTS

Comparative analysis of GlcN(acyl)PI in HeLa cell lines

In a previous study we reported on a HeLa S3 subline, denoted HeLa D, with unusually high levels of GlcN(acyl)PI (approx. 10^7 molecules per cell), that allowed us to perform compositional analyses of this GPI intermediate [6]. In the present study we analysed four HeLa cell lines to determine whether a high level of GlcN(acyl)PI is a common feature of HeLa cells or is a characteristic unique to the HeLa D subline. These cell lines were CCL-2, the original HeLa line; S3, a clone of CCL-2; D, derived from S3 [6]; and D3, a clone from D obtained by limiting dilution. The morphologies of CCL-2, S3 and D are quite different and might reflect their different sensitivity to growth inhibition by contact. Cells from the CCL-2 line appeared elongated, whereas HeLa D cells were rounded and did not form monolayers but rather grew in clusters (Figure 1). S3 cells showed a heterogeneous population of cells with both characteristics and clone D3 had an appearance similar to D.

The four HeLa cell lines were labelled for 3 days with...
Figure 2  GlcN(acyl)PI accumulates in HeLa D cells

Cells (10^6) from each HeLa cell line were incubated with 10 µCi of [3H]inositol for 3 days, the lipids extracted in chloroform/methanol/water (10:10:3, by vol.), dried and partitioned between butanol and water as outlined in the Experimental section. The butanol phases, containing approx. 500000 c.p.m., were run on TLC plates developed in chloroform/methanol/water (10:10:3, by vol.). The distribution of radioactivity shown was obtained by radioscanning the TLC plate.

Figure 3  Comparative analysis of [3H]inositol-labelled lipids from HeLa D and mutant E cells

HeLa D and lymphoma mutant E cells (10^6) were labelled with 10 µCi of [3H]inositol for 3 days. Labelled lipids were extracted from HeLa D and mutant E cells and samples containing equal amounts of protein (with approx. 500000 and approx. 300000 c.p.m. respectively) were analysed as described in the legend to Figure 2.

HeLa D cells produce higher GlcN(acyl)PI levels than mutant E

Lymphoma mutant E cells harbour a defect in Dol-P-Man synthase and as a consequence accumulate GlcN(acyl)PI [7]. We compared mutant E and HeLa D cells for their abilities to accumulate GlcN(acyl)PI. After [3H]inositol labelling to steady state, GlcN(acyl)PI was detected in E cells (Figure 3). After normalization for cell number, the smaller lymphoma cells incorporated only 10% of the added [3H]inositol label, in comparison with 50% in HeLa D. [3H]Inositol-labelled lipids from equivalent amounts of membranes from both cell lines were compared on TLC (Figure 3). HeLa D cells produced 4 times
more GlcN(acyl)PI than lymphoma E cells, and the ratio of GlcN(acyl)PI to PI in HeLa D cells was twice that in the mutant E cells.

GlcN(acyl)PI is the substrate for the first mannosyltransferase, which uses Dol-P-Man as the mannose donor. Two types of defect affecting Dol-P-Man can disrupt the normal mannosylation of GPIs and result in the potential for GlcN(acyl)PI accumulation: (1) total lack of Dol-P-Man synthase activity (as found in mutant E cells) or partial lack of Dol-P-Man synthase activity (as represented in CHO cell [17] and T-cell hybridoma mutants) [23], and (2) impaired Dol-P-Man utilization (as represented in Lee 35). In view of the above findings that HeLa D cells are the highest producers of GlcN(acyl)PI reported, the following experiments were designed to answer whether the defect in these cells involves Dol-P-Man synthesis or utilization.

**Dol-P-Man synthase activity is normal in HeLa D cells**

To compare Dol-P-Man synthase activity between HeLa cell lines, the synthesis of [3H]Dol-P-Man from [3H]GDP-Man and Dol-P was assayed, with mutant E as a negative control. As shown in Table 1, similar levels of Dol-P-Man synthase activity were observed in the HeLa cell lines and contrasted with the absence of activity in lymphoma mutant E. This result ruled out the possibility that HeLa D cells and the mutant E share the same defect. It is, however, noteworthy that these synthase activity measurements do not reflect the steady-state levels of Dol-P-Man or utilization of the Dol-P-Man pool.

**HeLa D cells can synthesize mannosylated-GPIs in the presence or absence of tunicamycin**

Cell mutants that are partly deficient in Dol-P-Man produce smaller amounts of mannosylated-GPIs and GPI-anchored proteins [17,23]. In these leaky Dol-P-Man synthase mutants, the levels of mannosylated-GPIs and GPI-anchored proteins can be boosted by the addition of tunicamycin [17,23]. Tunicamycin inhibits the synthesis of Glu-Man, GlcNac, PPDol, the precursor of N-glycans on proteins, and Dol-P-Man is then used only in the GPI pathway. For this reason mannone labelling of GPIs is routinely performed in the presence of tunicamycin.

We first investigated the levels of the mature mannosylated-GPIs (H6, H7 and H8) and Dol-P-Man in HeLa D cells and in the other HeLa cell lines in the presence of tunicamycin: after 1 h of [3H]Man labelling, H6, H7 and H8 were detected in HeLa D and D3 cells but the levels of 3H in the two latter intermediates and Dol-P-Man were lower than in the CCL-2 cells (Table 2). To rule out the possibility that HeLa D is a leaky Dol-P-Man synthase mutant, the amounts of endogenous Dol-P-Man available for GPI-mannosylation were decreased by omitting tunicamycin from the medium. In this condition, as expected, Dol-P-Man levels were decreased by 70% in CCL-2 cells but surprisingly were unchanged in S3 and S3-derived cells (Table 2). Furthermore, removal of tunicamycin caused no significant decrease in H6, H7 and H8 in HeLa D and D3 cells, indicating that in these cells Dol-P-Man was not limiting (Table 2). The relative levels of these Man,GPIs differed between the cell lines. H8 was the major [3H]Man-labelled Man,GPI species in CCL-2 cells, whereas similar levels of H6 and H8 were observed in the S3 and S3-derived cell lines (D and D3) (Table 2). Moreover, incorporation of [3H]Man into H7, H8 and Dol-P-Man was 2–4 times higher in CCL-2 cells than in the other cell lines (Table 2), although steady-state levels of H8 in HeLa D and CCL-2 measured by [3H]inositol labelling were comparable within the different cell lines (results not shown). The lack of an effect of tunicamycin on HeLa D GPIs together with the normal levels of Dol-P-Man synthase activity (Table 1) in these cells clearly indicates that they are not deficient in Dol-P-Man.

**HeLa D cells synthesize a Man9 lipid-linked oligosaccharide**

Lee 35 is a CHO mutant that expresses normal levels of Dol-P-Man but is unable to use this carrier as a mannose donor for the synthesis of lipid-linked oligosaccharides and presumably mannosylated GPIs [14]. This CHO mutant produces low levels of GPI-anchored proteins and apparently no mannosylated GPIs [14]. Because of an inactive Dol-P-Man pool, Lee 35 cells do not synthesize mature dolichol-linked oligosaccharides containing nine mannosyl residues but, as in the lymphoma mutant E cells [24], accumulate lipid-linked oligosaccharides having only five mannosyl residues [14]. One possible explanation for the fact that levels of Dol-P-Man did not rise in S3 and S3-derived cells (D and D3) when tunicamycin was added to the incubation medium (Table 2) is that in these cell lines Dol-P-Man is not used for the synthesis of lipid-linked oligosaccharides. To test this possibility, size analysis of the [3H]Man-labelled glycans released from dolichol-linked oligosaccharides from CCL-2 and D cells was performed by P4 exclusion chromatography. Glycans from both lines eluted with similar relative elution constants (Kd values of 0.24 and 0.23 respectively) (Figure 4), very close to the values reported for Glc2ManGlcNAc2 derived from many different cell types [25] but lower than those for the truncated species from mutant E cells ([25], and results not shown). The shoulder eluting

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**Table 1** Dol-P-Man synthase activity in HeLa cell lines and mutant E

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<tr>
<th>Cell line</th>
<th>Dol-P-Man (c.p.m./µg of protein)</th>
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<tr>
<td>CCL-2</td>
<td>34</td>
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<tr>
<td>S3</td>
<td>19</td>
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<tr>
<td>D</td>
<td>27</td>
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<tr>
<td>D3</td>
<td>20</td>
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<tr>
<td>E</td>
<td>0.2</td>
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**Table 2** Quantification of Man,GPIs and Dol-P-Man in HeLa cell lines

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Tu…</th>
<th>H6</th>
<th>H7</th>
<th>H8</th>
<th>Dol-P-Man</th>
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<tr>
<td>CCL-2</td>
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<td>2060</td>
<td>640</td>
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<td></td>
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increase Dol-P-Man levels in S3 and S3-derived cell lines. The fact that S3 and S3-derived cells make oligosaccharides (Figure 4), distinguishing this S3 subline from the synthesis of mannosylated GPIs (Table 2) and the formation of normal lipid-linked Dol-P Man in HeLa D cells is used in both the mannosylation of GPI-anchored proteins in the four HeLa cell lines, we labelled the cells with PI-PLC (approx. 15% respectively for the sample from CCL-2 cells and 58 and 158 respectively for the sample from HeLa D cells. Fractions of 0.45 ml were collected and taken for scintillation counting.

HeLa D cells express GPI-anchored proteins

The mammalian cell mutants noted above that are defective in the synthesis of mannosylated GPIs expressed decreased amounts of GPI-anchored proteins. To compare the profiles of GPI-anchored proteins in the four HeLa cell lines, we labelled the cells to a steady state with [3H]inositol and extracted the labelled proteins with Triton X-114. SDS/PAGE profiles from the four cell lines showed [3H]inositol-labelled proteins at 70, 40 and 20 kDa (Figure 5). PI-PLC treatment of the extracts shifted most of the radiolabel from the detergent phase to the aqueous phase, indicating that only GPI-anchored proteins are labelled with [3H]inositol. Residual label in the 70 kDa band after treatment with PI-PLC (approx. 15%) was probably due to fatty acid acylation of the inositol [26,27]. It is noteworthy, however, that the 70 kDa band from the S3 and S3-derived cells was heterogeneous and ran consistently more slowly than the 70 kDa band from CCL-2 (Figure 5). This apparent higher molecular mass did not seem to be caused by partial N-glycosylation because HeLa D cells synthesized a normal precursor of N-glycosylation (Figure 4), although it is possible that the glycan transfer to proteins is slow in these cells. To quantify this more rigorously, we focused on one GPI-anchored protein, decay accelerating factor (DAF), which is probably the most abundant GPI-anchored protein in these cells [28]. Both cell-surface DAF and total cell-associated DAF proteins were determined by immunoradiometric assay. When the amount of DAF in the supernatants was normalized to total cellular protein, no differences were found between the four HeLa cell lines in either cell-surface DAF (Table 3) or total DAF (results not shown).

Comparative pulse–chase of [3H]inositol in HeLa D and CCL-2 cells

To explore the mechanistic basis for the accumulation of GlcN(acyl)PI in HeLa D cells, we compared the rates of synthesis and metabolism of this species in pulse–chase experiments with [3H]inositol. For simplicity, results were interpreted in the context of a one-compartment model [Scheme 1 and eqns. (1)–(3)] that permitted rate constants to be estimated. The results shown in Figure 6 indicate that the rate constants for synthesis and metabolism of PI are similar for both cell lines. In contrast, the rate constant $k_2$ for GlcN(acyl)PI formation is approx. 40 times higher in HeLa D cells than in CCL-2, whereas no significant
HeLa D subline

![Pulse with \( ^{3}\text{H} \)inositol for 70 h](image)

HeLa D cells (a) or CCL-2 cells (c) (10^6 cells) in 1 ml of inositol-free medium were incubated with 1–10 µCi/ml \( ^{3}\text{H} \)inositol for the indicated times before extraction and analysis by TLC as outlined in the legend to figure 2. Total radioactivity in lipids was determined by scintillation counting of the butanol phase and the distribution of lipid species was determined by radioscanning of the TLC plates and scintillation counting of the silica beads. After 70 h of incubation with \( ^{3}\text{H} \)inositol, the HeLa D cells (b) or CCL-2 (d) were washed to remove radiolabel in the medium, returned to 10 ml of complete medium and extracted at the indicated times and analysed as in (a) and (c). The observed amounts of radioactivity in PI (B, _) and GlcN(acyl)PI (C, *), relative to the initial radioactivity of added inositol (A, 0), are plotted against time and the time courses for both panels with each line were fitted to eqns. (2) and (3) derived from the one-compartment model.

Data for HeLa D cells were fitted with the following values:

- \( k_A = 0 \)
- \( k_1 = 0.012 \text{ h}^{-1} \)
- \( k_2 = 0.017 \text{ h}^{-1} \)
- \( k_3 = 0.031 \text{ h}^{-1} \)
- \( k_4 = 0.022 \text{ h}^{-1} \)

Data for CCL-2 cells were fitted with the following values:

- \( k_A = 0 \)
- \( k_1 = 0.012 \text{ h}^{-1} \)
- \( k_2 = 0.010 \text{ h}^{-1} \)
- \( k_3 = 8 \times 10^{-5} \text{ h}^{-1} \)
- \( k_4 = 0.022 \text{ h}^{-1} \)

Difference in the rate constants of metabolism in these cell lines was observed.

**DISCUSSION**

In this paper we describe a HeLa S3 cell subline designated D that accumulates GlcN(acyl)PI while still producing mannosylated GPIs and GPI-anchored proteins. The HeLa D cells comprise a less adherent subpopulation of cells within the parental S3 line. We observed an inverse correlation between HeLa cell adherence and GlcN(acyl)PI synthesis. Adherence decreased progressively for the CCL-2, S3 and D lines. Reciprocally, D cells produced 4 times more GlcN(acyl)PI than S3 and almost 40 times more than CCL-2. It is noteworthy that HeLa D cells produce twice the levels of GlcN(acyl)PI observed in lymphoma mutant E, the mammalian cell line first reported to accumulate GlcN(acyl)PI (Figure 3).

This D cell line is unique because it does not fall in the same group of Dol-P-Man-deficient mutants previously found to accumulate GlcN(acyl)PI [7,13]. Dol-P-Man synthase activity and Dol-P-Man levels are normal in HeLa D cells (Table 1 and Table 2), and these characteristics clearly distinguish this cell line from the other mutants. A normal level of Dol-P-Man does not necessarily imply its normal utilization, as exemplified by Lec 35 cells [14], but HeLa D cells seem to use Dol-P-Man as a mannosyl donor in the synthesis of mannosylated-GPIs (Table 2) and lipid-linked oligosaccharides (Figure 4). Furthermore the levels of Man,GPIs in D cells were not affected by tunicamycin (Table 2), indicating that the amount of Dol-P-Man is not limiting the mannosylation of GlcN(acyl)PI. These results provide further evidence that HeLa D cells are not a leaky Dol-P-Man mutant with low levels of Dol-P-Man. The broader band of \( ^{3}\text{H} \)inositol-labelled GPI-anchored proteins at 70 kDa in all S3-derived cells (Figure 5), combined with the lack of effect of tunicamycin on Dol-P-Man levels in the S3 and S3-derived cell lines, raised the possibility that these cells had a glycosylation defect. Recent results on the topology of mannosylated-GPI biosynthesis have suggested that these reactions take place primarily on the cytoplasmic side of the ER [29]. In contrast, mannosylation of lipid-linked oligosaccharides is believed to involve Dol-P-Man on the luminal side of the ER [30]. If these topologies are assumed, the possible glycosylation...
defect could involve a faulty flippase that produces a normal GPI-mannosylation and defective N-glycan synthesis. This hypothesis was discarded because lipid-linked oligosaccharides in HeLa D cells contained the full nine-mannosyl glycan core (Figure 4). Normal utilization of Dol-P-Man in the GPI pathway (Table 2) and in the synthesis of this precursor of N-glycosylation of proteins (Figure 4) distinguishes HeLa D from Lec 35 cells.

HeLa D cells produce at least as much GPI-anchored protein as the cell lines from which they were derived (Figure 5 and Table 3). Yeast mutants that accumulate GPI intermediates without decreasing the levels of GPI-anchored proteins were also recently reported by Conzemian's group. Some of these mutants accumulated GPI anchor intermediates without effect on the levels of GPI-anchored proteins, whereas other mutants with close to normal amounts of GPI anchor intermediates showed low levels of GPI-anchored proteins [31].

The HeLa D cell phenotype could be explained if the cell population included two types of cells, in principle, one producing high levels of GlcN(acyl)PI and without mannosylated GPs or GPI-anchored proteins and the other with undetectable GlcN-(acyl)PI and normal amounts of Man,GPIs and GPI-anchored proteins. Clone D3 ruled out this possibility because it showed the same profiles of mannosylated GPs and GPI-anchored proteins as in the uncloned D cell line (Table 2 and Figure 5). Moreover, cell-surface DAF detected immunologically with monoclonal anti-DAF antibodies and revealed by a secondary fluorescent antibody showed that all HeLa D cells were uniformly stained and that the numbers of cells in the bright and fluorescent fields were the same (results not shown).

The high levels of GlcN(acyl)PI in HeLa D cells could result from an increased rate of its synthesis, a decreased rate of its metabolism or a combination of both effects. Because there is no increase in the levels of downstream intermediates, any increased synthesis cannot be accompanied by an increase in mannosylation, either because the mannosyltransferase is saturated or because GlcN(acyl)PI accumulates in a compartment inside or outside the ER where it cannot undergo rapid mannosylation. Kinetic modelling of the rates of approach to the steady state during pulse-chase experiments with [3H]inositol in CCL-2 and D cells indicated that the apparent rate constant for the conversion of PI to GlcN(acyl)PI in HeLa D cells was almost 40 times higher than that in CCL-2. In contrast, the combined rate constants for decay of GlcN(acyl)PI are comparable in both cell lines. A higher rate of synthesis of GlcN(acyl)PI in HeLa D cells must arise from an increase in the rate constant for the rate-limiting step preceding GlcN(acyl)PI formation. This could be GlcNAc addition to PI, deacetylation of GlcNACP or acylation of GlcNP. However, this step also could be the exit of GlcN(acyl)PI or an earlier precursor from the ER compartment where mannosylation is relatively rapid to an inert compartment where mannosylation is either slow or non-existent. Recent evidence supports such an exit to an inert compartment (A. Wonkanjorslip and T. Rosenberry, unpublished work). Although not yet identified, this compartment could be the plasma membrane.

Whatever the cause for the accumulation of GlcN(acyl)PI, the HeLa D subline has proved to be a useful tool by providing sufficient material for compositional analysis of GlcN(acyl)PI [6] and for allowing the demonstration that exogenous PI added to the cells can be metabolized to GlcN(acyl)PI [32]. We also believe that this HeLa subline will be useful for studies of the GlcN(acyl)PI mannosyltransferase and of the regulation of the GPI pathway.

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REFERENCES


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