Glucocorticoid-regulated Glycoprotein Maturation in Wild-type and Mutant Rat Cell Lines

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Abstract. Glucocorticoid hormones can regulate the posttranslational maturation of mouse mammary tumor virus (MTV) precursor polyproteins in M1.54, a stably infected rat hepatoma cell line. We have used complement-mediated cytolysis to recover variants of M1.54 that fail to express MTV cell surface glycoproteins in a hormone-regulated manner (Firestone, G. L., and K. R. Yamamoto, 1983, Mol. Cell. Biol., 3:149-160). One such clonal isolate, CR4, is similar to wild-type with respect to synthesis of MTV mRNAs, production of the MTV glycoprotein precursor (gPr74 env) and a glycosylated maturation product (gp51), and hormone-induced processing of two MTV phosphoproteins. In contrast, three viral cell surface glycoproteins (gp78, gp70, and gp32) and one extracellular species (gp70s), which derive from gPr74 env in glucocorticoid-treated wild-type cells, fail to appear in CR4. CR4 showed no apparent alterations in proliferation rate, cell shape, or expression of total functional mRNA and bulk glycoproteins. We conclude that the genetic lesion in CR4 defines a highly selective hormone-regulated glycoprotein maturation pathway that alters the fate of a restricted subset of precursor species.

P OLYPEPTIDES destined to be membrane associated or secreted undergo a series of specific processing and compartmentalization reactions during and after translation (21, 44, 50, 52, 57, 62, 63). Cell biological and genetic data (1, 11, 17, 48, 51, 52, 59) suggest that shuttle vesicles selectively transport polypeptides from the rough endoplasmic reticulum into the Golgi stacks, through the Golgi cisternae and to their final intra- or extracellular locations, apparently subjecting the polypeptides to an ordered array of modifications. For example, different Golgi cisternae appear to contain specific glycosyltransferases that modify glycoprotein substrates during passage through individual compartments (7, 8, 10, 21, 44). In addition, fatty acid modification of certain polypeptide backbones appears to occur at specific loci within the Golgi region (7, 9, 55). Different proteins with similar final destinations appear to use separate transport pathways (15, 52, 58), and certain pathways seem to include branch points that specify alternative processing fates (3, 14, 19, 31). Given the complexity, generality, and overall importance of these events, it would not be surprising if they were exploited for regulation of the relative levels of specific final products. Indeed, secretion of ACTH (19, 32) and parathormone (6) may be modulated by cAMP, although little is known of the mechanisms of these phenomena.

Glucocorticoids appear to regulate the posttranslational maturation of mouse mammary tumor virus (MTV) glycoproteins and phosphoproteins in M1.54, an MTV-infected rat hepatoma cell line containing 10 stably integrated proviruses (13, 14). For example, the predominant viral glycoprotein produced by control cultures is an extracellular 51,000 Mr species (gp51s), whereas hormone-treated cells produce two extracellular forms (gp51s and gp70s), and three cell surface species (gp78, gp70, and gp32). In addition, dexamethasone facilitates maturation of two new viral phosphoproteins (p24 and p35), derived by proteolytic cleavage from their common phosphorylated polypeptide (14, 24). Finally, the hormonal signal has a third effect on MTV gene expression, stimulating the rate of MTV RNA synthesis by increasing selectively the efficiency of MTV promoter utilization (5, 18, 38-40, 43, 46, 53, 60, 61, 64). Thus, glucocorticoids govern the absolute levels as well as the maturation of MTV proteins in M1.54. The present study, together with previous evidence (13), suggests that the three glucocorticoid-regulated pathways are genetically distinct.

The normal cellular targets for the hormone-controlled posttranslational processes have not been identified; however, the MTV polypeptides provide sensitive and perhaps general substrates for identifying and characterizing the regulated pathways. For example, we have selected and analyzed M1.54 variants defective in the maturation of certain MTV glycoprotein species in the presence of hormone. One such mutant line, CR4, expresses MTV RNA normally but fails to produce mature cell surface viral glycoproteins (13). In this report, we define molecular features of hormone-regulated glycoprotein maturation by comparing the expres-
tion and processing of MTV glycoproteins in the parent and mutant cell lines.

Materials and Methods

Materials

L-[35S]Methionine (1,000 Ci/mmol) and d-[2-3H]mannose (16 Ci/mmol) were obtained from Amersham Corp. (Arlington Heights, IL); [9,10-3H(N)]-palmitic acid was from New England Nuclear (Boston, MA); dexamethasone was from Sigma Chemical Co. (St. Louis, MO); and agaroase derivatized with castor bean lectin (Ricin)-l20 was from P-L Biochemicals, Inc. (Milwaukee, WI). All antisera were generous gifts of L. J. T. Young and R. D. Cardiff (Department of Pathology, University of California, Davis); endo-β-N-acetylglucosaminidase H (endo H) was a gift of Regis Kelly (Department of Biochemistry and Biophysics, University of California, San Francisco). All other reagents were of highest available purity.

Cells and Method of Culture

M1.54 is a cloned line of MTV-infected rat hepatoma tissue culture (HTC) cells (47); selection of its complement-resistant derivative CR4 has been described previously (3). Cells were cultured in DME supplemented with 5% horse serum as described (14). Where indicated, dexamethasone was present in the medium throughout both incubation periods. Hormone-treated cultures were preincubated with dexamethasone (1 μM and emission wavelengths of 405 and 465-540 nm, respectively, in a Model H Turner fluorimeter. Under these conditions, a reading of one fluorescence unit equals 10^4 cells.

RNA Isolation and In Vitro Translation

Total RNA was isolated from M1.54 and CR4 cells exposed to 1 μM dexamethasone, translated in vitro in a rabbit reticulocyte lysate (41), and [35S]methionine-labeled proteins were analyzed by one-dimensional gel electrophoresis (35) as described previously (13).

Results

Analysis of Mutant CR4 Proliferation, Cell Shape, and Functional mRNA

Using complement-mediated cytolysis, the CR4 mutant line was isolated on the basis of its inefficient expression of cell surface MTV polypeptides in the presence of dexamethasone (13). Conceivably, gross differences in cell growth rate or shape could give rise to this phenotype. Glucocorticoids have been shown to affect cell proliferation, in particular producing a nearly complete inhibition of cell growth in certain hepatoma lines (20). As shown in the graph in Fig. 1, M1.54 and CR4 each display doubling times of ~25 h in the absence of hormone and both show only a modest (18%) decrease in growth rate in the absence of dexamethasone. Similarly, light microscopic comparisons of CR4 and M1.54 cultures propagated 6 d in the presence or absence of hormone (Fig. 1; four images), revealed no systematic differences in cell shape; independent of hormone treatment, each cell line was epithelial-like and somewhat flattened with polygonal structures. Thus, in contrast to a previous report (23), we concluded that cell shape per se does not appear to account for the defective localization of cell surface MTV polypeptides in CR4.

To survey the expression of functional mRNA in M1.54 and CR4, total cellular RNA was isolated from hormone-induced cells, translated in vitro in a rabbit reticulocyte ly-
Figure 1. Proliferation and shape of dexamethasone-treated and untreated MI.54 and CR4 cells. (Left) MI.54 and CR4 cells were plated in the presence or absence of 1 μM dexamethasone and the DNA content was analyzed by a diaminobenzoic acid fluorescence assay through a 6-d time course. Using a standard curve, DNA-mediated fluorescence was monitored at the indicated times and converted to cell number per cm². (Right) At the 6-d time point, cells were examined by light microscopy. Bar, 50 μm.

Figure 2. Two-dimensional gel electrophoresis of in vitro translation products from MI.54 and CR4 cells. The cell-free translation system was programmed with 10 μg of total RNA isolated from MI.54 (top) or CR4 (bottom) that had been propagated in the presence of 1 μM dexamethasone. Samples (2 μl) were analyzed by two-dimensional electrophoresis with 80% pH 5–8 and 20% pH 3.5–10 ampholines in the isoelectric focusing (IEF) dimension. The molecular mass standards are β-galactosidase (120,000 Mₐ), BSA (68,000 Mₐ), ovalbumin (43,000 Mₐ), aldolase (40,000 Mₐ), and chymotrypsigen (25,000 Mₐ). The arrows denote the 42,000 Mₐ (50-kD secreted glycoprotein) and 22,000 Mₐ (α₁-acid glycoprotein) in vitro translation products induced by dexamethasone.

M1.54 -DEX → CR4 -DEX △○
M1.54 +DEX ← CR4 +DEX △△

M1.54 -DEX
M1.54 +DEX

CR4 -DEX
CR4 +DEX

sate (41), and the translation products analyzed by twodimensional gel electrophoresis (35). As shown in Fig. 2, MI.54 and CR4 RNAs yielded nearly identical patterns of in vitro translation products, suggesting that CR4 expresses normally the vast majority of its functional mRNA; consistent with this view, the overall rates of protein synthesis in hormone-induced and uninduced CR4 and MI.54 are indistinguishable as assessed by the incorporation of [³⁵S]methionine during 5-min pulse labeling (data not shown). Interestingly, comparison of the gels from Fig. 2 with those of translation products from untreated cultures revealed two major hormone-inducible species in both lines, α₁-acid glycoprotein (22,000 Mₐ) and a 50,000 Mₐ secreted glycoprotein (43,000 Mₐ primary product). Thus, the lesion in CR4 appears to affect the expression of a restricted subset of gene products while general competence for glucocorticoid responsiveness is retained.
Expression of MTV Glycoproteins in M1.54 and CR4

To examine MTV glycoprotein maturation, M1.54 and CR4 cultures exposed to 1 μM dexamethasone for 24 h were pulse-labeled with [2-3H]mannose followed by additional incubation in the presence of excess unlabeled mannose. At various times, [2-3H]mannose-labeled viral glycoproteins were immunoprecipitated with anti-MTV antibodies, fractionated in SDS polyacrylamide gels, and visualized by fluorography. As shown in Fig. 3 (lanes E–H), dexamethasone-treated M1.54 initially synthesizes a 74,000 M₉ precursor glycoprotein (designated gPr74e°V) that is subsequently matured into four major species (gp70, gp70, gp51, and gp32); several other minor or transiently expressed viral glycoproteins are also produced. Hormone-treated CR4 also produces gPr74e°V (upper panel, lane J), but its subsequent processing is dramatically different (lanes J–L); gp78, gp70, and gp32 expression is reduced or undetectable in CR4, whereas several other gPr74e°V-derived species, most notably gp51, are produced competently. We have suggested previously that specific portions of the MTV protein maturation pathways are dexamethasone inducible. To compare the hormone-regulated species in M1.54 with those deficient in CR4, uninduced cultures of M1.54 were pulse-labeled with [2-3H]mannose, followed by subsequent incubation with excess mannose, immunoprecipitation of MTV glycoproteins, and fractionation by electrophoresis. As shown in Fig. 3 (lanes A–D), untreated cells synthesize gPr74e°V, but produce little detectable [2-3H]mannose-labeled gp78, gp70, and gp32. Thus, the production of gPr74e°V-derived glycoproteins in control M1.54 (lanes A–D) is strikingly similar to that in hormone-treated CR4 (lanes I–L), suggesting that the genetic lesion in CR4 specifically blocks the hormonal regulation of glycoprotein maturation.

This notion was tested further by analyzing the extracellular viral glycoproteins. We established previously that at steady state, hormone-treated M1.54 produce two free extracellular MTV polypeptides (70,000 M₉ and 51,000 M₉) in approximately equimolar amounts, while control cells secrete predominantly (>95%) the 51,000 M₉ species (14). Fig. 3 shows glycoproteins pulse-labeled with [2-3H]man-
Figure 4. Cell surface-associated and intracellular MTV glycoproteins. Hormone-treated (24 h) M1.54 cells were either radiolabeled at the cell surface by lactoperoxidase-mediated $^{[25]}$I-iodination (lanes A–E) or metabolically labeled for 24 h with 300 μCi $^{[2-3H]}$mannose (lanes F–J). Cell extracts were immunoprecipitated with anti-MTV antibodies (lanes A and F), monospecific anti-gp51 antibodies (lanes B and G), monospecific anti-p28 antibodies (lanes C and H), monospecific anti-p28 antibodies (lanes D and I), or preimmune serum (lanes E and J). Immunoprecipitated polypeptides were fractionated by SDS PAGE and visualized by autoradiography or fluorography.

Identification of Cell Surface MTV Glycoproteins

The expression in CR4 of a specific subset of gPr74-derived maturation products suggests that the CR4 mutation may uncouple two distinct pathways for MTV glycoprotein maturation, one of which yields cellular gp78, gp70, and gp32 in the presence of hormone. To examine the relationships of these hormone-regulated glycoproteins and to determine those species located at the cell surface, plasma membrane proteins accessible on intact cells were labeled by lactoperoxidase-catalyzed $^{[25]}$I-iodination (Fig. 4, lanes A–E), while a parallel culture was metabolically labeled with $^{[2-3H]}$mannose (Fig. 4, lanes F–J). Immunoprecipitation with anti-MTV antibodies (lanes A and F) revealed that the iodinated 78,000 M₉, 70,000 M₉, 51,000 M₉, and 32,000 M₉, cell surface species co-migrate in SDS gels with the predominant viral glycoproteins gp78, pg70, gp51, and gp32, respectively, whereas several of the minor MTV glycoproteins are not detected at the cell surface. Monospecific anti-gp51 antibodies recognized both iodinated and $^{[2-3H]}$mannose-labeled forms of the 78,000 M₉, 70,000 M₉, and 51,000 M₉, species (Fig. 4, lanes B and G) while monospecific anti-gp32 antibodies recognize a cell surface 32,000 M₉ glycoprotein (Fig. 4, lanes C and H) in addition to the two stable polyproteins (78,000 M₉ and 70,000 M₉ species). A minor $^{[125]}$I-labeled species migrating at 20,000 M₉ was also detected by anti-gp32 antibodies and may represent a specific proteolytic fragment of gp32. In contrast, none of the iodinated cell surface or $^{[2-3H]}$mannose-labeled viral polypeptides were detected with preimmune or antiphosphoprotein sera.

These results demonstrate that the cell surface MTV polypeptides are glycosylated, and that gp32 and gp51 are immunologically distinct, while gp78 and gp70 appear to be polyproteins containing all or portions of the polypeptide backbones for both gp51 and gp32. In fact, sequence analysis of the GR virus env gene revealed that gp51 is the amino-
terminal portion of the polyprotein while gp32 is the carboxy-terminal domain and contains the transmembrane hydrophobic region (45). Interestingly, gp51 is associated with the cell surface despite its lack of a membrane anchor; conceivably gp51 may be associated with another primary integral membrane protein during transport to the cell surface. Consistent with this mechanism, we have shown that after multiple freeze-thaw at -100°C of isolated microsomes, which elutes luminal components of vesicles, gp51 remains associated with the membrane fraction (Haffar, O. K., C. P. Edwards, and G. L. Firestone, manuscript submitted for publication). Finally, the cell surface viral glycoproteins that fail to appear in CR4 (gp78, gp70, and gp32) have in common the polypeptide backbone for gp32 suggesting a crucial maturation or sorting domain may be contained within its structure. Thus, defective posttranslational maturation of the MTV glycosylated precursor appears to account for the absence of gp78, gp70, and gp32 from the CR4 cell surface. Since M1.54 and its derivatives contain 10 nontandemly integrated MTV proviruses (13), all of which seem transcriptionally active (Harris, C., and K. R. Yamamoto, unpublished data), the CR4 defect is likely to reflect a lesion in a cellular gene, rather than a mutation in the env gene itself or elsewhere within MTV sequences.

Endo H Sensitivity of MTV Glycoproteins

In general, glycoprotein modification and maturation include addition of "high mannose" moieties in the rough endoplasmic reticulum which are further processed during transit through the Golgi stacks into "complex" oligosaccharide sidechains (21, 27, 28). Endo H, which specifically hydrolyzes the high-mannose species, was used to examine oligosaccharide maturation of [35S]methionine-labeled MTV glycoproteins from dexamethasone-treated cells. As shown in Fig. 5, both externalized species, gp51s and gp70s (whose expression in CR4 is reduced several fold), appear to contain a heterogeneous mixture of endo H-resistant and sensitive carbohydrate side chains. Moreover, the pattern of endo H sensitivity is indistinguishable in M1.54 and CR4, implying that the altered maturation of particular MTV glycoproteins in CR4 is not due simply to a global defect in oligosaccharide processing.

Endo H treatment of [35S]methionine-labeled MTV glycoproteins immunoprecipitated from both M1.54 and CR4 yielded a 62,000 M₉, gP74env derivative; a parallel experiment with [2-3H]mannose-labeled glycoproteins revealed that all of the [2-3H]mannose in gP74env is removed (data not shown). In contrast, the [2-3H]mannose radiolabel in the membrane-associated gp78, gp70, gp51, and gp32 products remained attached, as expected for complex oligosaccharide side chains (data not shown). Finally, the glycosylated MTV precursors produced in M1.54 and CR4 migrate indistinguishably in two-dimensional gels (data not shown), suggesting that they are identical.

Attachment of [3H]Palmitate to MTV Glycoproteins

For certain plasma membrane proteins, covalently attached fatty acid residues appear to "anchor" the protein within the membrane (35); in the case of the vesicular stomatitis virus G protein, this lipid modification likely occurs during transport through the cis Golgi region soon after the acquisition of endo H resistance (7). When M1.54 and CR4 cultures were incubated for 24 h with [3H]palmitate in the presence of 150 μM dexamethasone, only the hormone-inducible gp78, gp70, and gp32 species became labeled (Fig. 6, right panel, lane C); in particular, no palmitate-labeled viral proteins were detected in either control (lane E) or hormone-treated (lane G) CR4. Labeling of parallel cultures with [35S]methionine (Fig. 6, left panel), showed that CR4 failed to express the polypeptide backbones for gp78, gp70, and gp32, rather than producing aberrant species unable to acquire the lipid modification.
Figure 6. Analysis of \([\text{H}]\)palmitic acid–labeled and \([\text{S}]\)methionine–labeled viral proteins. (Left) M1.54 (lanes A and B) and CR4 cells (lanes C and D) were exposed to 1 \(\mu\)M dexamethasone for 24 h and radiolabeled with 25 \(\mu\)Ci \([\text{S}]\)methionine during the last 4 h of hormone treatment. Proteins immunoprecipitated with anti-MTV antibodies (lanes A and C) or preimmune serum (lanes B and D) were fractionated in SDS polyacrylamide gels and visualized by fluorography. Pr74 denotes the position of two distinct MTV polyprotein precursors encoding the glycosylated and phosphorylated viral proteins; p35 and p24 are mature MTV phosphoproteins (see references 14 and 22).

(Right) M1.54 (lanes A–D) and CR4 (lanes E–H) cells were labeled with 0.5 mCi \([\text{H}]\)palmitic acid for 24 h in the presence (lanes C, D, G, and H) or absence (lanes A, B, E, and F) of 1 \(\mu\)M dexamethasone, and extracts were immunoprecipitated either with anti-MTV antibodies (lanes A, C, E, and G) or with preimmune serum (lanes B, D, F, and H); radiolabeled MTV proteins were fractionated by SDS gel electrophoresis and analyzed by fluorography. Molecular weight standards for both panels were as in Fig. 2.

These results revealed that palmitate modification is a biochemical feature common to the cell surface MTV glycoproteins not expressed in CR4 cells. It is clear, however, that palmitate modification itself is neither hormone-dependent in M1.54 nor defective in CR4, since the specific activity of \([\text{H}]\)palmitate incorporated into TCA-precipitable material from dexamethasone-treated and untreated M1.54 and CR4 are similar (Table I). Table I also shows that \([\text{H}]\)fucose and \([2,3\text{H}]\)mannose incorporation levels in the two lines are equivalent; moreover, hormone-treated and control M1.54 and CR4 produce similar \([\text{H}]\)palmitate-modified and \([2,3\text{H}]\)mannose–labeled cellular glycoproteins (Haffar, O. K., and G. L. Firestone, unpublished data), establishing further that bulk glycoprotein expression in CR4 is unaffected.

**Discussion**

A variety of sorting mechanisms operate upon glycoproteins to effect their maturation and transit to membrane-associated or extracellular locations (17, 21). For example, in active or carrier-mediated pathways, vesicle or organelle-associated "sorting receptors" have been postulated to bind and segre-

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<th>Cell line</th>
<th>Dexamethasone</th>
<th>([\text{H}])Palmitate</th>
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* Cells were incubated with or without 1 \(\mu\)M dexamethasone for 24 h in the presence of 1 mCi \([\text{H}]\)palmitate, detergent solubilized, and the level of radioactive lipid incorporated into protein was determined by precipitation with 10% TCA. Specific incorporation is expressed as cpm/mg of precipitated protein.

† The incubation and assay conditions were the same as described in previous footnote, except that the cells were incubated with 200 \(\mu\)Ci of \([2,3\text{H}]\)mannose.

§ The incubation and assay conditions were the same as described in the first footnote, except that the cells were incubated with 200 \(\mu\)Ci of \([\text{H}]\)fucose.
gate selectively the appropriate glycoprotein substrates (2, 25), presumably by recognizing features of the polypeptide backbone or of the oligosaccharide side chains, as with the mannose-6-phosphate receptors involved in targeting lysosomal hydrolases (10, 56). Previous work of others suggests that some of these processes may be amenable to genetic dissection in cultured cells (57, 59). For example, Kabat and coworkers have isolated mutants defective in processing Friend murine leukemia virus or Friend spleen focus-forming virus envelope glycoproteins (16, 49), and Zilberstein et al. (65) have recovered variants that fail to produce cell surface-associated and lipid-modified vesicular stomatitis virus gene products. The novel feature of our results is the identification of a defect in a hormone-regulated posttranslational process. We conclude that the MTV glycoprotein precursor can be partitioned differentially into either a glucocorticoid-inducible or a constitutive trafficking pathway in cultured rat hepatoma cells. This finding demonstrates that global physiological regulators such as steroid hormones can effect acute changes in the products expressed from specific genes even after their transcription.

The ability of hormone-treated CR4 and control cultures of M1.54 to survive in the presence of MTV-specific antiserum and complement reflects the absence of high levels of cell surface-associated gp78, gp70, and gp32; only in hormone-treated M1.54 does gPr74 env proceed efficiently through the maturation and sorting pathway that results in the membrane localization of these species. Thus, dexamethasone appears to mediate an increase in a particular subset of transport or targeting activities or to stimulate the entry of gPr74 env into a pre-existing sorting route. It may be relevant in this regard that the cell surface glycoproteins not expressed in CR4 contain the polypeptide backbone for gp32 (the COOH-terminal domain of gPr74 env), perhaps indicating that a sorting signal resides within that structure.

In any case, the genetic lesion in CR4 displays a highly restricted phenotype, blocking selectively the hormonal induction of a glycoprotein maturation pathway that operates upon a specific subset of target precursors. Notably, the glucocorticoid receptor appears normal in CR4, and we have demonstrated that several hormone-responsive genes are regulated competently (13). Moreover, the gPr74 env species produced in hormone-treated and control M1.54 and CR4 are indistinguishable with respect to molecular weight, migration in two-dimensional gels, glycosidase sensitivity, and lectin affinity. In addition, bulk glycoprotein processing and compartmentalization appear normal in CR4, in contrast, for example, to the global alterations in intracellular protein traffic seen in the yeast secretory mutants defined by Schechman and coworkers (33, 34, 54). Similarly, control and hormone-treated M1.54 and CR4 are indistinguishable in the production of total palmitate-derivatized polypeptides (data not shown). Since the membrane-destined MTV glycoproteins normally produced in hormone-treated M1.54 and absent in CR4 are all palmitate modified, it seems likely that the hormone-regulated branch point resides at or before palmitate addition.

The regulatory mechanism by which the glucocorticoid receptor alters glycoprotein processing and compartmentalization is unknown. The putative hormone-regulated factors identified by the CR4 mutation might function directly in the processing of specific glycoproteins, or may operate as intracellular sorting receptors (25), analogous to the actions of egsyn and β2-microglobulin, which are essential for membrane association of β-glucuronidase (37) and HLA antigens (29, 36), respectively. Alternatively, the regulated factors might facilitate membrane vesicle flow into particular intracellular compartments, or maintain localization of specific target proteins subsequent to vesicle fusion (10, 25). In any case, inhibitors of RNA synthesis prevent the hormone-regulated glycoprotein maturation events (Haffar, O. K., and G. L. Firestone, manuscript submitted for publication). Thus, one simple interpretation is that glucocorticoids may regulate the transcription of genes encoding certain receptors or processing enzymes, and that the expression of one or more of these genes is defective in CR4.

Using MTV glycoproteins as targets for genetic selection and as intracellular biochemical probes, we have been able to detect and characterize a regulated posttranslational glycoprotein expression pathway without indications a priori of its existence, its normal cellular target proteins, or its biological significance. Conceivably, a pathway of this type could efficiently redirect the maturation and targeting fates of even large existing pools of glycoproteins simply by altering the synthesis or degradation of a single factor. This could facilitate subtle or dramatic transitions in specific glycoprotein localization, such as might be involved, for example, in cell membrane remodeling during development and differentiation. Of course, the precise nature of the regulated events and the regulatory signals themselves are likely to be cell specific. Thus, cAMP appears to control the externalization of mature ACTH in pituitary cells (19, 32). Through further characterization of CR4 and other genetic variants, it should be possible to identify cellular proteins whose maturation and targeting are controlled by glucocorticoids and to elucidate the normal biological function of this phenomenon.

This report is dedicated to the memory of Edward C. Heath, who supervised G. L. Firestone's doctoral research from 1975-1980 and whose guidance and friendship will always be remembered. We thank the colleagues in our laboratories for their helpful comments and suggestions during the course of this study. We also thank Aileen Kim in the Department of Physiology-Anatomy, University of California, Berkeley, for her superb typing and preparation of this manuscript; and John Underhill in the Cancer Research Laboratory, University of California, Berkeley, for his excellent photography.

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References

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plemented groups required for post-translational events in the yeast secretory

storage, proteolytic processing, and secretion on stimulation.

29. Krangel, M. S., H. T. Orr, and J. L. Strominger. 1979. Assembly and


12. Firestone, G. L. 1983. The role of protein glycosylation in the compart-

16. Fitting, T., M. Ruta, and D. Kabat. 1981. Mutant cells that abnormally

15. Fitting, T., and D. Kabat. 1982. Evidence for a glycoprotein "signal" in-

14017.


29. Krangel, M. S., H. T. Orr, and J. L. Strominger. 1979. Assembly and

30. Laemmli, U. K. 1970. Cleavage of structural proteins during the assem-

34. Novick, P., C. Field, and R. Schekman. 1980. Identification of 23 com-


38. Payvar, F. P., D. DeFranco, G. L., Firestone, B. Edgar, O. Wrangle, S.


43. Ponta, H., N. Kennedy, P. Skroch, N. E. Hynes, and B. Groner. 1985. Hormonal response region in the mouse mammary tumor virus long terminal repeat can be dissociated from the proviral promoter and has enhancerproper-


46. Renkawitz, R., G. Shutz, D. von der Ahe, and M. Beato. 1984. Se-


52. Saraste, J., and E. Kuismanen. 1984. Pre- and post-Golgi vacuoles oper-


59. Tartakoff, A. M. 1983. Mutations that influence the secretory path in ani-


63. Wold, P. 1981. In vivo chemical modification of proteins (posttransla-


65. Zilberstein, A., M. D. Saider, M. Porter, and H. F. Lodish. 1980. Mu-


68. Madin-Darby canine kidney cells, d.


72. Ponta, H., N. Kennedy, P. Skroch, N. E. Hynes, and B. Groner. 1985. Hormonal response region in the mouse mammary tumor virus long terminal repeat can be dissociated from the proviral promoter and has enhancerproper-


75. Firestone et al. Regulation of Glycoprotein Trafficking 2331