Compartment-Restricted Biotinylation Reveals Novel Features of Prion Protein Metabolism in Vivo

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Proteins are often made in more than one form, with alternate versions sometimes residing in different cellular compartments than the primary species. The mammalian prion protein (PrP), a cell surface GPI-anchored protein, is a particularly noteworthy example for which minor cytosolic and transmembrane forms have been implicated in disease pathogenesis. To study these minor species, we used a selective labeling strategy in which spatially restricted expression of a biotinylating enzyme was combined with asymmetric engineering of the cognate acceptor sequence into PrP. Using this method, we could show that even wild-type PrP generates small amounts of the CtmPrP transmembrane form. Selective detection of CtmPrP allowed us to reveal its N-terminal processing, long half-life, residence in both intracellular and cell surface locations, and eventual degradation in the lysosome. Surprisingly, some human disease-causing mutants in PrP selectively stabilized CtmPrP, revealing a previously unanticipated mechanism of CtmPrP up-regulation that may contribute to disease. Thus, spatiotemporal tagging has uncovered novel aspects of normal and mutant PrP metabolism and should be readily applicable to the analysis of minor topologic isoforms of other proteins.

INTRODUCTION

A characteristic feature of complex organisms is their ability to markedly diversify the protein products generated from a comparatively small number of genetic elements. Through the use of alternative promoter usage, alternative splicing, translational regulation, modification, processing, regulated degradation, and trafficking, a single gene may generate any of numerous potential protein products in a highly context dependent manner. While the major protein product is typically the most extensively studied, it is increasingly appreciated that minor or transiently generated alternative protein species often have critical functional roles.

For example, many proteins are primarily made and reside in an inactive form, with the active form represented by minor or transiently produced products. Hence, the minor species of key factors like steroid hormone receptors (Yudt and Cidlowski, 2002), membrane-bound transcription factors (Sannerud and Annaert, 2009), or caspasas (Pop and Salvesen, 2009) are in fact the primary functional product.

Even seemingly artificial minor truncated versions of Glucocorticoid receptor resulting from internal translation initiation seem to be physiologically relevant functional products in vivo (Lu and Cidlowski, 2005; Lu et al., 2007). Thus, understanding the nuances and regulation of complex organism physiology will require an understanding of not only how proteomic complexity is generated, but the functional roles of the various major and minor products.

However, studying the alternative minor species of a protein often poses several technical obstacles depending on its comparative abundance and difference from the major species. In the simplest case, the minor species can be differentiated on the basis of a unique sequence element (e.g., generated by an isoform-specific exon) that can be immunologically detected with high specificity and sensitivity. However, this is not always possible. For example, alternative forms generated via facultative translocation or trafficking can generate minor variants whose main distinguishing feature is its alternative cellular location.

Indeed, proteins with targeting signals for the ER, mitochondria, peroxisomes, and chloroplasts can potentially reside in the cytosol as a minor (or ‘eclipsed’) nontranslocated form (reviewed by Regev-Rudzki and Pines, 2007). While this is sometimes apparently attributable to errors (Anandatheethavarada et al., 2003; Drisaldi et al., 2003; Rane et al., 2004; Levine et al., 2005), numerous examples have been described where the noncompartmentalized form has a function (see Yoge et al., 2010, for an especially compelling recent example). This species can be generated by alternative splice variants that lack the targeting signal (Tong et al., 2003), alternative translation initiation sites downstream of the targeting signal (Land and Rouault, 1998; Goulet et al., 2004), modifications or processing close to the targeting signal (Addya et al., 1997; Robin et al., 2002; Colombo et al.,...)

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inefficiencies in the targeting or translocation process (Stein et al., 1994; Sass et al., 2001; Shafer et al., 2005; Regev-Rudzki et al., 2005; Naamati et al., 2009; Matthews et al., 2010). Studying these minor, differentially compartmentalized, and often transiently generated species has been challenging. Yet, they are nonetheless important. In cases where they represent mistakes, their quality control and degradation is likely to help maintain protein homeostasis in the cytosol. In cases where they have functional relevance, delineating their function requires their selective identification and analysis.

One example where alternative species are thought to be important is mammalian Prion protein (PrP). While the function of this widely expressed cell surface GPI-anchored glycoprotein remains to be clearly elucidated (Westergaard et al., 2007; Aguzzi et al., 2006), it is best known for causing various fatal neurodegenerative diseases when misfolded in certain ways (Prusiner, 1998; Aguzzi and Calella, 2009). Recently, several cell biological studies have focused on the possibility that mislocalization of PrP can be detrimental to the cell and perhaps be a contributing factor in the pathogenesis of at least some of the associated diseases (reviewed in Chakrabarti et al., 2009). One source of PrP mislocalization stems from a slight inefficiency in the function of its N-terminal signal sequence (Kim et al., 2002; Rane et al., 2004; Levine et al., 2005). This typically results in ~10% failed translocation into the ER (Rane et al., 2004; Levine et al., 2005), generating a minor cytosolic form of PrP termed cyPrP (Ma and Lindquist, 2002; Drisaldi et al., 2003; Rane et al., 2004). However, signal sequence inefficiency can have another outcome. If, after targeting to the translocon, the signal sequence fails to initiate translocation of the N terminus, an internal hydrophobic domain (HD, at residues ~112–135) can engage the translocon and direct membrane insertion of PrP (Kim et al., 2001; Kim et al., 2002; Kim and Hegde, 2002; see Supplemental Figure S1). This transmembrane form, termed CtmPrP, has the N terminus in the cytosol and C terminus in the ER lumen (Hegde et al., 1998). Importantly, mutations that increase hydrophobicity of the hydrophobic domain result in increased CtmPrP production (Hegde et al., 1998; Kim et al., 2001; Stewart and Harris, 2001, 2003).

Studies in transgenic mice manipulating PrP translocation in several ways have illustrated that increased generation of either cyPrP or CtmPrP can cause neurodegeneration (Hegde et al., 1998, 1999; Ma et al., 2002; Stewart et al., 2005; Rane et al., 2008). Of note, several human disease mutations are in the central hydrophobic domain that, based on in vitro assays, result in increased CtmPrP (Hegde et al., 1998; Stewart and Harris, 2001; Kim and Hegde, 2002; Rane et al., 2010). One of these, A117V, has also been modeled in transgenic mice and shown to cause a late-onset neurodegeneration (Hegde et al., 1999; Yang et al., 2009; Rane et al., 2010) and increased CtmPrP (Hegde et al., 1999; Rane et al., 2010). As predicted from the above description of how cyPrP and CtmPrP are generated, improving signal sequence efficiency resulted in reduced generation of these forms in vitro (Kim and Hegde, 2002) and in transgenic mice (Rane et al., 2010), and rescued mice from neurodegeneration (Rane et al., 2010). Thus, CtmPrP and cyPrP, while minor species of PrP, may be of considerable importance to the pathogenesis of at least some diseases. However, these forms, particularly CtmPrP, are especially difficult to detect and follow in biochemical analyses, and this has severely limited insights into its biosynthesis, trafficking, and metabolism.

The difficulty in detecting and analyzing the minor cyPrP and CtmPrP forms is primarily an issue of signal to noise. In the case of CtmPrP, the polypeptide is identical (or nearly so) to the major fully translocated species: both are glycosylated and GPI-anchored, migrate similarly on SDS-PAGE, and are composed of the same polypeptide sequence (Hegde et al., 1998; Stewart et al., 2001; Kim and Hegde, 2002; Stewart and Harris, 2003). Thus, selective antibody detection is difficult. While conformation-specific antibodies are a theoretical possibility, this poses substantial technical hurdles given that the specific conformations are not easy to generate in high amounts as a source of antigens. Thus, to study CtmPrP, a method to selectively mark this minor species is needed. Here we have explored the possibility of using compartment-restricted and site-specific biotinylation as a method for selective tagging of CtmPrP. We characterize this method for use in mammalian cells and use it to delineate basic but key features of CtmPrP biosynthesis and metabolism. Illustrating the power of this approach, we show that, unexpectedly, CtmPrP metabolism is selectively perturbed by disease-associated mutants in PrP previously unknown to affect PrP topology.

**MATERIALS AND METHODS**

**DNA Constructs**

Wild-type Hamster PrP and the various derivatives from it [ASS-PrP, SA-PrP, PrP(G123P), PrP(G123P), PrP(AV3), PrP(G123P)] have been described (Hegde et al., 1998, 1999; Ma et al., 2002; Rane et al., 2004; Chakrabarti and Hegde, 2009). Wild-type Human PrP and its various disease-associated point mutants have been described (Ashok and Hegde, 2009). In each case, the BioTag or an HA epitope is at residues 109–112. PrP-A recognizes all mammalian species of PrP and recognizes Hamster PrP, but not SA-PrP. These were used interchangeably with indistinguishable results, except in experiments with SA-PrP, in which case PrP-A was used. The TRAPs antibody has been described (Fons et al., 2003).

**Antibodies**

Anti-PrP antibodies used in this study were either PrP-A, a rabbit polyclonal against the PrP N terminus from residues 23–38 (Ashek and Hegde, 2008; Rane et al., 2008), or 3F4 (Covance, Princeton, NJ), a mouse monoclonal whose epitope is at residues 109–112. PrP-A recognizes all mammalian species of PrP and recognizes Hamster PrP, but not SA-PrP. These were used interchangeably with indistinguishable results, except in experiments with SA-PrP, in which case PrP-A was used. The TRAPs antibody has been described (Fons et al., 2003).

**Cell Culture**

Unless otherwise indicated, all experiments were performed in Neuro 2a (N2a) cells obtained from ATCC. Key results were confirmed to be reproducible in HeLa, HEK293, and HT1080 cells (unpublished results). To facilitate much of the routine characterization, N2a cells stably expressing BirA were produced by isolating and selecting for clones from single cells using standard methods. However, transient cotransfection with BirA produced identical results and stable expression was not essential. In instances where BirA was transiently transfected, it occupied between one-sixth and one-fifth of the total DNA. Transfection was with either Lipofectamine 2000 or Effectene with similar results. Cells were assayed between 24 and 48 h after transfection. In nearly all experiments, a small amount (typically one-fifth of the total amount of the DNA) of a CFP or GFP expression plasmid was cotransfected to verify that efficiency of transfection was at least 70%. Although not shown, this was often used as an expression control. Experiments were typically performed on cells at ~70–90% confluence. Pulse-chase studies were performed on cells in six-well dishes as described (Rane et al., 2004; Ashok and Hegde, 2008; Ashok and Hegde, 2009). Although adding exogenous biotin to the media was not absolutely necessary, the experiments here contained 10 μM biotin in the culture media that was typically added at the time of transfection.
**Compartment-Restricted Tagging Method**

**In Vitro Biotinylation Assays**

In vitro transcription, translation in reticulocyte lysate, and translocation using canine pancreatic rough microsomes has been described (Kim et al., 2002; Fons et al., 2003; Sharma et al., 2010). The protease protection assay for topology was as before (Kim et al., 2002). For biotinylation, translation reactions were performed as above but included 1 nM PrP-A antibody (Ashok and Hegde, 2008) or 3F4 antibody (Covance). EndoH and PNGase treatments and analysis of PrP solubility in non-denaturing detergents was as before (Rane et al., 2004; Ashok and Hegde, 2008; Ashok and Hegde, 2009). BirA was expressed in E. coli BL21(DE3) pLYsS cells with IPTG (1 mM) induction at midlog phase growth. Expression was for 3 h, after which the cells were harvested in 300 mM NaCl, 50 mM Tris, pH 8 and lyed by sonication. The His-tagged BirA was purified by immobilized Ni$^{2+}$ or Co$^{2+}$ affinity chromatography (using Chelating-Sepharose from GE), dialyzed against PBS and stored in aliquots at −80°C. SDS-PAGE was on 12% Tris-Tricine gels, except the experiments in Figure 6, which were on 15% Tris-Tricine gels.

**Quantification**

 Autoradiographs were quantified by phosphorimaging on a Typhoon system, while images for the figures were produced by exposure of the same gel to film and scanning in Photoshop. Comparative quantification of western blots were by comparing samples to serial dilutions of standards that were run on the same blot.

**RESULTS AND DISCUSSION**

**Experimental Strategy**

To selectively identify and monitor $^{13}$CmPrP, we exploited its unique topology relative to the other PrP forms: cytosolically exposed N-terminal domain with C-terminal domain in the ER lumen. Because the C-terminal domain has acceptor sites for N-linked glycosylation, its residence in the lumen can be readily monitored. To generate a similar topologic and cytosolically exposed N terminus, we took advantage of orthogonal biotinylation of an artificial short sequence by the bacterial BirA biotin ligase (Schatz, 1993). This enzyme, which covalently attaches biotin to its substrate proteins, has been found by screening and refinement to biotinylate a short artificial acceptor sequence of 13 residues (Schatz, 1993; Beckett et al., 1999; hereafter termed the BioTag; see Figure 1A). This BioTag, which contains an acceptor lysine for biotinylation, is efficiently modified by BirA, is not found in any natural proteins, and is not recognized by mammalian biotinylating enzymes (Schatz, 1993; Beckett et al., 1999; Howarth et al., 2005; Kulman et al., 2007). This makes the BirA-BioTag system orthogonal, allowing the generation of a highly specific and easily detectable modification.

We therefore reasoned that PrP BioTagged within its N-terminal domain could allow selective $^{13}$CmPrP detection (Figure 1B). When this Biotagged PrP (PrP$^{Bio}$) is expressed in cells containing BirA in the cytosol, only $^{13}$CmPrP should be both biotinylated and glycosylated. The majority of PrP would be fully translocated into the ER lumen cotranslationally, thereby escaping biotinylation (but not glycosylation). By contrast, cytosolic PrP would be biotinylated but not glycosylated, and $^{13}$CmPrP would be neither glycosylated nor biotinylated. Thus, using two topologically restricted modifications allows, in principle, the four potential forms of PrP to be discriminated (Figure 1B). In practice however, any potential deficiencies in the modification reactions could make the lack of a modification difficult to interpret with certainty, especially when dealing with low abundance species. For this reason, and because of its clear physiological importance, we focused on $^{13}$CmPrP, whose double modification can be interpreted with a high degree of confidence.

Our overall strategy was to develop and validate this compartment-restricted tagging strategy in vitro and in cells using variants of PrP with well-defined and uniform (or nearly uniform) topology. Once validated, we sought to then apply the method to wild-type PrP so that basic but critical unanswered questions about $^{13}$CmPrP could be addressed. And finally, we investigated whether inherited mutants in human PrP that cause neurodegeneration might influence $^{13}$CmPrP metabolism.

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**Miscellaneous Biochemistry**

Immobilized ConA was from GE Biosciences (Piscataway, NJ) and was used at 10 µl per 1 ml sample. Incubation was for 2 h at 4°C in IP buffer as for the Avdin pulldowns. Immunoprecipitations were as before and used the anti-
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The 13-residue BioTag, with a flanking Alanine on either side, was inserted into PrP at codon 50, between the flexible N-terminal 27 residues and the octarepeat region. Insertion of a short tag in this region is not expected to have any effect on PrP based on earlier studies inserting either an epitope tag or GFP (Negro et al., 2001; Lorenz et al., 2002; Chakrabarti and Hegde, 2009; Ashok and Hegde, 2009). Thus, throughout this study, all constructs were modified at this site and the modified constructs indicated by a superscript ‘Bio’ (or ‘HA’ for the HA epitope tag) in the name. Two versions of PrP that are uniformly (or very nearly so) in the CtmPrP or SA-PrP Bio capture were tested in an in vitro system (Figure 2A).

As characterized before (Chakrabarti and Hegde, 2009), SA-PrP(G123P)Bio translated in rabbit reticulocyte lysate in the presence of canine pancreatic rough microsomes was inserted efficiently into the membrane as judged by its glycosylation, cosedimentation with microsomes, protease accessibility of its N terminus, and protease protection of its C terminus. By contrast, Prl-PrP(G123P)Bio (containing the highly efficient N-terminal 27 residues and the octarepeat region. Insertion of a short tag in this region is not expected to have any effect on PrP based on earlier studies inserting either an epitope tag or GFP (Negro et al., 2001; Lorenz et al., 2002; Chakrabarti and Hegde, 2009; Ashok and Hegde, 2009). Thus, throughout this study, all constructs were modified at this site and the modified constructs indicated by a superscript ‘Bio’ (or ‘HA’ for the HA epitope tag) in the name. Two versions of PrP that are uniformly (or very nearly so) in the CtmPrP or fully translocated topologic forms were BioTagged and tested in an in vitro system (Figure 2A).

When these translation products were incubated with recombinant BirA, SA-PrP(Bio) translated in rabbit reticulocyte lysate in the presence of canine pancreatic rough microsomes was inserted efficiently into the membrane as judged by its glycosylation, cosedimentation with microsomes, protease accessibility of its N terminus, and protease protection of its C terminus. By contrast, Prl-PrP(G123P)Bio (containing the highly efficient N-terminal 27 residues and the octarepeat region. Insertion of a short tag in this region is not expected to have any effect on PrP based on earlier studies inserting either an epitope tag or GFP (Negro et al., 2001; Lorenz et al., 2002; Chakrabarti and Hegde, 2009; Ashok and Hegde, 2009). Thus, throughout this study, all constructs were modified at this site and the modified constructs indicated by a superscript ‘Bio’ (or ‘HA’ for the HA epitope tag) in the name. Two versions of PrP that are uniformly (or very nearly so) in the CtmPrP or fully translocated topologic forms were BioTagged and tested in an in vitro system (Figure 2A).

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Figure 1. Experimental strategy for compartment-restricted biotinylation. (A) Diagram of PrP (top) and SA-PrP (bottom) indicating key elements: SS – signal sequence; HD – hydrophobic domain; CHO – glycosylation site; GPI – signal for glycosylphosphatidylinositol anchor attachment; SA – signal anchor (from Asialoglycoprotein receptor). In addition, the BioTag and control HA sequences are shown at their site of insertion. The Lysine that becomes biotinylated is in red. SA-PrP lacks a N-terminal signal sequence and is instead targeted to and inserted via the SA domain. The flanking charges ensure its type II orientation, resulting in the C terminus being translocated into the ER lumen. Numbering is for Human PrP. (B) Diagram of the four potential topologic forms that can be achieved by PrP, with the glycan sites indicated in green, and the BioTag in red. On introduction of BirA to the cytosol, cyPrP and CtmPrP become biotinylated (‘B’). A chart summarizing the modifications is shown.

Analysis of Compartment-Restricted Biotinylation in Vivo

To evaluate the suitability of using biotinylation as a means to detect CtmPrP in vivo, we coexpressed SA-PrP(Bio) with BirA in N2a cells and analyzed total cell lysates for biotinylated products by blotting with Avidin-HRP. In preliminary experiments, we initially confirmed that cells expressing BirA without a BioTagged construct, or a BioTagged construct without BirA, showed the same weakly staining bands as observed in GFP-transfected control cells (data not shown). This indicated that BirA has no detectable targets endogenous to mammalian cells, and that the BioTag is not modified by endogenous enzymes. Similar results were seen in multiple cell types (data not shown), where the main endogenous products detected by Avidin-HRP were two high molecular weight bands characterized before as mitochondrial biotin-containing enzymes (Howarth et al., 2005).

When SA-PrP(Bio) was introduced into BirA-expressing cells, it was clearly visualized on blots with both anti-PrP antibodies and Avidin-HRP (Figure 3A). SA-PrP biotinylation was entirely dependent on the BioTag, because SA-PrP lacking the BioTag was not detected by Avidin-HRP despite expression at high levels as judged by its detection with anti-PrP. The presence of biotin in the media maximized expression at high levels as judged by its detection with anti-PrP. The presence of biotin in the media maximized expression at high levels as judged by its detection with anti-PrP. The presence of biotin in the media maximized expression at high levels as judged by its detection with anti-PrP. The presence of biotin in the media maximized expression at high levels as judged by its detection with anti-PrP.
glycosylated and unglycosylated PrP verified the glycan specificity of ConA binding under the conditions used in this study (Supplemental Figure S2). Furthermore, blotting of lysates before and after incubation with immobilized Avidin illustrated that most of the SA-PrPBio was biotinylated (Figure 3C). The residual amount remaining in the unbound fraction was likely due to inefficiency in Avidin binding using batch-pulldown. Thus, a construct expected to be in the CtmPrP form can be site-specifically and efficiently biotinylated in vivo with cytosolic BirA. That this biotinylated species is also glycosylated confirms its CtmPrP topology.

**Figure 3.** Biotinylation assay for PrP topology in vivo. (A) SA-PrP and ΔSS-PrP lacking or containing the BioTag were transfected into BirA-expressing N2a cells and analyzed 24 h later. Total cell lysates or ConA-bound products were separated by SDS-PAGE, blotted, and probed with either anti-PrP antibodies (α-PrP) or Avidin-HRP as indicated. Note that low levels of endogenous PrP are visualized at ~35 kDa in the ΔSS-PrP samples probed with α-PrP. (B) SA-PrPBio was transfected into N2a cells expressing or lacking BirA and cultured in the indicated concentrations of biotin for 24 h. Total cell lysates were analyzed for biotinylated products by blotting with Avidin-HRP. (C) N2a cells expressing BirA were transfected with either a GFP-expressing plasmid (control), SA-PrP, or SA-PrPBio. Total cell lysates were incubated with immobilized Avidin to pull-down the biotinylated proteins. Samples of the total lysate, unbound fraction, and bound fraction were analyzed by immunoblotting with anti-PrP. (D) The indicated constructs were transfected into N2a cells expressing either cytosolic BirA or SS-BirA-KDEL (ER-BirA). Total cell lysates were analyzed by blotting and probing for either total PrP (α-PrP) or biotinylated products (with Avidin-HRP). ‘2HA’ indicates a construct containing two tandem HA tags.
As with SA-PrP<sub>Bio</sub>, a cytosolic version of PrP lacking its ER targeting signal (ASS-PrP<sub>Bio</sub>) was also biotinylated by BirA in a BioTag-dependent manner (Figure 3A). This is noteworthy because nontargeted cytosolic PrP is rapidly degraded, and the ability to detect its biotinylation illustrates that even short-lived species that are likely chaperone-associated can be tagged by BirA.

We also examined the consequences of directing BirA to the ER lumen with an N-terminal signal sequence and C-terminal KDEL sequence (SS-BirA-KDEL). In principle, this should preclude biotinylation of a cytosolically exposed BioTag, while permitting biotinylation of a BioTag in the ER lumen. As expected, biotinylation of SA-PrP<sub>Bio</sub> was clearly decreased (Figure 3D). In contrast, wild-type PrP<sub>Bio</sub>, the majority of which is expected to be translocated completely into the ER lumen, was biotinylated to a greater extent by ER-targeted BirA than by cytosolic BirA (Figure 3D). While generally consistent with expectations, biotinylated and glycosylated SA-PrP<sub>Bio</sub><sub>G123P</sub> was still detectable in cells expressing SS-BirA-KDEL. This could be due to either residual SS-BirA-KDEL in the cytosol (i.e., due to translocation inefficiency), small amounts of SA-PrP<sub>Bio</sub><sub>G123P</sub> being fully translocated into the ER lumen, or post-lysis biotinylation.

We favor the first explanation for two reasons. First, sensitive reporter assays have previously shown that signal-containing proteins are never translocated with complete efficiency, leaving at least a few percent in the cytosol (Levine et al., 2005). Second, mixing experiments of cell lysates from BirA-expressing cells with lysates from PrP<sub>Bio</sub>-expressing cells showed no detectable post-lysis biotinylation under either the denaturing or nondenaturing lysis conditions we used (Supplemental Figure S3). This is presumably because upon washing and lysis of the cells, the concentrations of BirA, substrate, and biotin are reduced many orders of magnitude below their cellular levels, making the reaction very inefficient even if nondenaturing conditions are used. Thus, we conclude that while the majority of SS-BirA-KDEL is translocated and resides in the ER lumen [where it is apparently quite active, as observed before (Mize et al., 2008)], sufficient amounts of it are mislocalized to the cytosol where it can partially biotinylate a cytosolically exposed BioTag. Because SS-BirA-KDEL does not represent an absolute compartment-restricted enzyme, we focused the remainder of our efforts on using cytosolic BirA.

Detection of CtmPrP Generated by Wild-Type PrP

The SA-PrP model protein illustrated that biotinylation combined with glycosylation provides a rigorous means to detect CtmPrP since each modification is strictly restricted to different compartments. We sought to apply this method to unambiguously address whether CtmPrP is a normal product of wild type PrP biosynthesis. Although CtmPrP is generated from wild-type PrP in vitro, it is a minor product that some have argued represents inefficiencies or mistakes of an in vitro reaction comprising heterologous components (such as reticulocyte lysate and microsomes from pancreas). Whether CtmPrP is generated in vivo has been debatable because the assays for its detection have limited sensitivity and rely on either cumbersome biochemical fractionation (such as isolation of microsomes) or relatively crude limited protease digestion assays.

To address this issue, PrP<sub>Bio</sub> was coexpressed with cytosolic BirA, and the products were analyzed by various methods. PrP, whether it contains or lacks the BioTag, is expressed as a heterogeneously glycosylated set of products, with a small amount of unglycosylated material (Figure 4A). Detection of the biotinylated species with Avidin-HRP showed several BioTag-dependent PrP species (Figure 4A). The smallest of these products comigrated with unglycosylated PrP and was verified as such by its lack of binding to immobilized ConA. Furthermore, this band was the most efficiently biotinylated PrP product relative to its abundance in the total PrP blot, suggesting a high degree of cytosolic exposure. These observations suggest this species represents cyPrP, although it may also include CtmPrP that failed to receive N-linked glycans. Among glycosylated PrP species, a small proportion (~5–10%) was detected with Avidin-HRP. Most of these products could be pulled down with immobilized ConA and visualized by Avidin-HRP, confirming its dual modification with biotin and N-glycan, consistent with the CtmPrP topology.

In addition to detection of steady state levels of biotinylated PrP species, we could also selectively recover these forms from pulse-labeled cells (Figure 4B). As with the steady state analysis, capture of pulse-labeled PrP with immobilized Avidin was dependent on the BioTag. The assignment of bands as glycosylated and unglycosylated products was again confirmed using ConA binding (Figure 4C). Comparing the total pulse-labeled PrP<sub>Bio</sub> (recovered using anti-PrP immunoprecipitation) to that recovered with immobilized Avidin showed that unglycosylated PrP<sub>Bio</sub> was biotinylated to higher specific activity than glycosylated PrP<sub>Bio</sub>. Relative to glycosylated SA-PrP<sub>Bio</sub>, glycosylated PrP<sub>Bio</sub> was biotinylated to ~10-fold lower levels. Given that SA-PrP<sub>Bio</sub> is uniformly in the CtmPrP topology, this indicates that only a small proportion of glycosylated PrP<sub>Bio</sub> is represented by CtmPrP (~10% in this experiment). Although the precise proportion of PrP<sub>Bio</sub> in the CtmPrP topology varied depending on cell culture conditions, cell type, expression level, labeling time, and other parameters yet to be delineated, it was always many-fold lower than SA-PrP and was always a detectable but minor proportion of total synthesized PrP. These observations are consistent with unglycosylated PrP<sub>Bio</sub> representing primarily cytosolic species, while the glycosylated PrP<sub>Bio</sub> represents a mixture of fully translocated species with minor amounts of CtmPrP.

To further validate this conclusion, we tested whether biotinylation could be decreased by two independent manipulations that each reduce CtmPrP production. In the first strategy, we replaced the PrP signal sequence with that from Prolactin (Prl). Earlier studies have shown that both CtmPrP and cyPrP are dependent on slight inefficiency of the native PrP signal sequence, and that both can be reduced (although not completely eliminated) by the more efficient Prl signal (Kim and Hegde, 2002; Rane et al., 2004; Kang et al., 2006; Rane et al., 2010). Indeed, relative to PrP<sub>Bio</sub>, Prl-PrP<sub>Bio</sub> was reduced in the amount of both glycosylated and unglycosylated biotinylated PrP products recovered from pulse-labeled cells (Figure 4D). When total cell lysates were analyzed by blotting with Avidin-HRP, Prl-PrP<sub>Bio</sub> again showed a substantially reduced biotinylated population compared with PrP<sub>Bio</sub> despite equal or higher overall expression levels (Figure 4E).

In the second strategy, we analyzed PrP(G123P)<sub>Bio</sub>, which reduces CtmPrP levels by altering the potential membrane spanning region (e.g., Figure 2, C and D and Hegde et al., 1998). As with Prl-PrP<sub>Bio</sub>, we found that biotinylation of PrP(G123P)<sub>Bio</sub> was reduced approximately threefold, very close to the limits of reliable detection above background. The reduced biotinylation seen with PrP(G123P)<sub>Bio</sub> is noteworthy because it indicates that little if any biotinylation is occurring during cotranslational protein targeting. If this were the primary source of biotinylated product, PrP<sub>Bio</sub> and PrP(G123P)<sub>Bio</sub> would have been biotinylated equally since
failed to be glycosylated. Our ability to detect CtmPrP generated by wild-type PrP was pulse-labeled with [35S]methionine for 30 min. Total cell lysates were then immunoprecipitated with anti-PrP antibodies or pulled down (after its emergence from the ribosome (Krieg et al.)) with identical kinetics. The biotinylation assay reveals CtmPrP generated by wild-type PrP. (A) Wild-type Human PrP constructs lacking or containing the BioTag were transfected into BirA-expressing N2a cells and analyzed 24 h later. Total cell lysates or ConA-bound products were separated by SDS-PAGE, blotted, and probed with either anti-PrP antibodies or pulled down with immobilized Avidin. (D) An experiment as in panel B was performed on the indicated constructs. (E) BirA-expressing N2a cells were transfected with the indicated constructs, and total cell lysates were analyzed by blotting with anti-PrP or Avidin HRP. The positions of the unglycosylated and core-glycosylated products are indicated. The double-asterisk indicates complex glycosylated PrP species. (F) An experiment as in panel B was performed on the indicated constructs.

Three additional points argue against cotranslational biotinylation. First, SRP binds to the signal sequence immediately after its emergence from the ribosome (Krieg et al., 1986; Kurzchalia et al., 1986), even before the BioTag is exposed, thereby initiating the targeting process before the BioTag is available for modification. Second, earlier work in vitro and in vivo suggests that after the signal emerges from the ribosome, targeting occurs before even ~30 additional residues are synthesized (Jungnickel and Rapoport, 1995; Goder et al., 2000). During this period in PrP translation, the BioTag would reside inside the ribosomal tunnel, where it cannot be accessed by the large ~40 kDa BirA enzyme. Third, even if targeting were delayed, steric hindrance by the ribosome and SRP would likely preclude efficient access to the nascent PrP by BirA. For these reasons, both experimental and theoretical, we believe that biotinylation represents a post-translational event that is reporting on the final topologic location of the BioTag.

Thus, on the basis of dual modification by two topologically restricted enzymes (BirA and Oligosaccharyl transferase) and the results of various mutants that influence PrP topology [SA-PrP, ΔSS-PrP, Prl-PrP, and PrP(G123P)], we conclude that biotinylated and glycosylated PrPbio represents CtmPrP. While unglycosylated biotinylated PrP clearly includes cyPrP, it may well be a mixture with CtmPrP that failed to be glycosylated. Our ability to detect CtmPrP generated by wild-type PrP with high specificity at both steady state and by pulse-labeling paved the way to an analysis of this pathologically important form of PrP.

**Characterization of CtmPrP Trafficking and Metabolism**

With this assay for CtmPrP in hand, we addressed several basic, but still unresolved, questions regarding its normal biosynthesis and metabolism. Earlier studies using highly CtmPrP-favoring mutants [such SA-PrP or L9R-PrP(AV3)] have come to conflicting conclusions about the metabolism and cellular location of CtmPrP, ranging from ER, Golgi, or various other parts of the secretory pathway (Hegde et al., 1998; Stewart et al., 2001; Stewart and Harris, 2005; Chakrabarti and Hegde, 2009). However, these mutants are rather substantially modified, and it is unclear whether conclusions derived from them are entirely applicable. Whether CtmPrP generated from wild-type PrP ever leaves the ER, and if it does, whether it samples the cell surface, is unknown.

We found that the majority of biotinylated glycosylated PrPbio (i.e., CtmPrP) was sensitive to endoglycosidase H (EndoH) digestion (Figure 5A), indicating that most glycans on CtmPrP have not been processed by Golgi enzymes. However, a small amount of more slowly migrating bands was consistently observed, and these species were resistant to EndoH digestion, only being deglycosylated with PNGase F. The same species was also found to be accessible to extracellular trypsin, while the majority of biotinylated PrP was inaccessible (Figure 5B). When total PrP in the same samples was analyzed by blotting for PrP, the majority of PrP was...
downs, we estimated the half-life of CtmPrP as being before further analysis to ensure complete recovery of all experiments were harvested and solubilized in boiling SDS some of the earlier conflicting reports on its properties. Thus, A. B. Emerman et al. found it difficult to reliably detect a biotinylated complex 2xCHO 0xCHO Avidin pulldowns

Figure 5. Analysis of CtmPrP metabolism. (A) Cells expressing wild-type PrPBi and BirA were either left untreated or digested with extracellular trypsin. After washing and collecting the intact cells, they were harvested and either analyzed directly, or after capture of biotinylated products with immobilized Avidin. The blot was probed with anti-PrP antibodies. (B) Total lysates from cells expressing PrPBi and BirA were either left untreated or digested with Endoglycosidase H or PNGase F as indicated. The samples were then analyzed directly or after capture of biotinylated products with immobilized Avidin. The blot was probed with anti-PrP antibodies to visualize PrP, or anti–TRAPα, an ER-resident glycoprotein that serves as a control for the glycosidase digestions. (C) Cells expressing PrPBi and BirA were pulse labeled with [35S]methionine for 30 min and chased with unlabeled media for the indicated times. Lysates harvested at each time point were divided in half and immunoprecipitated with anti-PrP or pulled down with immobilized Avidin. The percent of glycosylated biotinylated PrP species remaining at each time point was quantified by phosphoimaging and indicated above the respective lanes. (D) Cells expressing PrPBi and BirA were treated for 16 h with the indicated compounds: 10 μM MG132, 250 nM Bafilomycin A1, 5 μM Lactacystin, or 0.2 mM Leupeptin. DMSO served as a vehicle control. Total cell lysates were subjected to Avidin pulldowns and the biotinylated products visualized with anti-PrP immunoblotting.

EndoH resistant and cell surface exposed (Figure 5, A and B). These observations indicate that in contrast to the major population of PrP, which is at the cell surface, CtmPrP is primarily intracellular, possibly in the ER as judged by its EndoH-sensitive glycans. However, at least some CtmPrP clearly leaves the ER and goes to the cell surface as evidenced by an EndoH-resistant population accessible to extracellular trypsin.

We also noticed in the course of our studies that CtmPrP was more difficult to solubilize in nondenaturing detergents than non-Ctm forms (unpublished observations; see also Figure 7A below). While insolubility could mean aggregation, we do not believe this to be the case because aggregates of a membrane protein would be unlikely to traffic through the secretary pathway and be modified on its glycans en route. Rather, it seems that two modes of membrane anchoring on the same molecule (via a transmembrane segment and GPI anchor) make it more difficult to solubilize. Regardless of the reason, this observation provides a further point of distinction between CtmPrP and the major PrP species. It also emphasizes the need to use efficient solubilization methods (e.g., SDS) to avoid losing subpopulations of PrP in any biochemical analyses. Indeed, standard lysis buffer conditions are insufficient to recover CtmPrP, perhaps explaining some of the earlier conflicting reports on its properties. Thus, unless specifically used for fractionation purposes, all of our experiments were harvested and solubilized in boiling SDS before further analysis to ensure complete recovery of all PrP species.

Using pulse-chase analysis combined with Avidin pulldowns, we estimated the half-life of CtmPrP as being ~5–7 h (Figure 5C). Consistent with the steady state analysis, most PrP synthesized during the pulse-labeling subsequently matured to heterogeneous complex glycosylated forms, after which it was degraded. By contrast, the majority of CtmPrP remained core-glycosylated. By the pulse-chase method, we found it difficult to reliably detect a biotinylated complex glycosylated species due to a low signal from this diffuse band relative to background. However, the steady state analysis does indicate that some proportion of CtmPrP does undergo complex glycans modification and traffic to the cell surface (Figure 5, A and B). It is worth noting that although most of CtmPrP remains core-glycosylated, its residence in the ER cannot be assumed because glycoproteins can be trafficked through the Golgi without necessarily obtaining further glycan modifications. At this point, we do not know the full itinerary of CtmPrP trafficking, although it does appear to sample much of the secretary and endocytic pathways given its eventual degradation in the lysosome (see below).

We used inhibitors to assess the normal degradation pathway used by CtmPrP. We focused on the ratio of glycosylated biotinylated products (i.e., CtmPrP) to unglycosylated biotinylated PrP (i.e., primarily cyPrP), because the latter is well established to be degraded by the proteasome. Relative to untreated cells (where the CtmPrP and cyPrP bands are roughly equal at steady state), proteasome inhibition (with either MG132 or Lactacystin) clearly results in cyPrP levels exceeding CtmPrP (Figure 5D). By contrast, lysosomal inhibition with either Bafilomycin A1 or Leupeptin resulted in increased CtmPrP relative to cyPrP (Figure 5D). These results indicate that while cyPrP is primarily degraded by a proteasome-dependent pathway, CtmPrP is primarily degraded in lysosomes.

And finally, we wished to assess the status of signal sequence processing on CtmPrP. Although a seemingly esoteric point, this issue is important for two reasons. First, it is quite unusual for a signal sequence containing protein to be generated in a topology where the N terminus is cytosolic. This necessarily means that the polypeptide segment downstream of the signal sequence has failed to be translocated into the ER, a fate inconsistent with signal sequence function. The step at which this fails in vivo has been unclear. If it fails at the step of targeting to or initial gating of the
translocon, the signal sequence will remain unprocessed. However, slippage of the polypeptide after initiating translocation could result in signal cleavage if it has transiently accessed the ER lumen where the signal peptidase active site resides. The second reason for assessing this is that because the signal sequence is hydrophobic, its lack of processing on CmPrP would generate a domain that may be prone to aggregation and/or inappropriate interactions that could play a role in cellular dysfunction caused by CmPrP. While analysis of a CmPrP-favoring mutant showed lack of signal sequence processing (Stewart et al., 2001), this construct contains a mutation in the signal sequence itself that may have affected the results.

To address this issue, we initially isolated CmPrP on the basis of its glycosylation and biotinylation and analyzed its migration relative to signal cleaved and uncleaved standards (data not shown). However, interpretation of this result proved more complicated than expected because of uncertainties about the effect of GPI anchor processing and deglycosylation (which converts Asparagines to Aspartates) on migration. To avoid these complications, we introduced a TEV cleavage site in the N-terminal domain downstream of the BioTag. After binding the glycoproteins (which would include CmPrP) to immobilized ConA, the N-terminal domain of PrPBio/TEV was liberated by digestion with TEV and the biotinylated N terminus (which necessarily is from CmPrP) was purified with immobilized Avidin (Figure 6A). This short product was then compared with signal-containing and signal-lacking versions of the fragment generated in vitro.

Figure 6. The signal sequence has been processed on CmPrP. (A) Experimental strategy to isolate the N-terminal domain selectively from CmPrP. Diagram of the TEV site inserted into PrP Bio. The two Methionines preceding the TEV site are to allow labeling; the cleavage site is indicated with an arrowhead. The fate of the four topologic forms of PrP upon ConA pulldown, TEV elution, and Avidin pulldown are shown. The final fragment recovered should necessarily be from CmPrP because it was selected for both glycosylation and biotinylation. (B) Characterization of the ConA pulldown and TEV cleavage steps. Pulse-labeled cell lysates expressing PrP Bio without or with the TEV site were analyzed directly (total lysate), after ConA pulldown, after TEV cleavage of ConA-selected products (ConA → TEV), after anti-PrP immunoprecipitation (α-PrP IP), and after TEV cleavage of the PrP IP products (α-PrP → TEV). The products were visualized by autoradiography. Note that no digestion occurs in the absence of the TEV site, while all of the PrP products are digested efficiently by TEV. PrP is barely visible in the ConA-selected products, which contains many endogenous cellular glycoproteins. Analysis was on 15% Tricine gels to maximize separation of small products. (C) Samples treated as in panel B were further selected by Avidin pulldowns and visualized by autoradiography. Markers for PrP from residue 1 up to the TEV site (+SS) or residue 23 up to the TEV site (−SS) were produced by in vitro translation and analyzed on the same gel. Note that an unidentified background translation product was seen in both samples (asterisks). The source of this product is unknown, although the nuclease digested translation system likely contains numerous mRNA fragments, some of which may be translated to generate small truncated products not normally visualized at the bottom of a gel. Analysis of this experiment was on 15% Tricine gels.
We first confirmed that TEV would specifically and efficiently cleave immobilized PrP on beads. To do this, PrP was immunoprecipitated with immobilized antibodies and after washing, incubated with TEV protease. PrPBio/TEV, but not PrPBio, was efficiently digested to yield the expected cleavage products (Figure 6B). Parallel samples on immobilized ConA also showed evidence that PrPBio/TEV was being digested similarly (Figure 6B). When the released products from the ConA digestion were further purified on immobilized Avidin, an ~7-kDa band was specifically recovered in a TEV protease and PrPBio/TEV dependent manner (Figure 6C). Comparison of this product to in vitro translation products of signal-containing and signal-lacking fragments showed that the TEV-cleaved product from in vivo generated CmPrP comigrates with the signal-lacking marker fragment (Figure 6C). Thus, CmPrP generated in vivo appears to have its signal sequence removed, suggesting that it is generated by slippage of the N terminus out of the translocon at a step after the polypeptide has transiently accessed the ER lumen. The absence of this hydrophobic peptide may explain why CmPrP is not recognized as a misfolded substrate for degradation by ER-associated degradation, while earlier studies generating CmPrP by mutating the signal sequence (which precluded its processing) came to different conclusions (Stewart et al., 2001).

A Disease-Causing PrP Mutant that Alters CmPrP Metabolism

An increased proportion of PrP is generated in the CmPrP form when residues in the central hydrophobic domain (HD) are mutated to more hydrophobic residues. This includes artificial mutants such as PrP(AV3), in which three alanines are changed to valines, as well as several naturally occurring disease-associated human PrP mutants (such as A117V; Hegde et al., 1998; Hegde et al., 1999; Kim et al., 2001; Stewart and Harris, 2003). Based on analyses of such HD
mutants in transgenic mice, increased levels of $\text{CtmPrP}$ have been linked to the development of neurodegeneration (Hegde et al., 1998; Hegde et al., 1999; Stewart et al., 2005; Yang et al., 2009; Rane et al., 2010). It is therefore thought that $\text{CtmPrP}$ may contribute to at least some genetic human disease caused by such PrP mutations in the central HD. However, numerous PrP mutants lie outside the HD and do not increase $\text{CtmPrP}$ production at the ER as assayed in vitro (Stewart and Hogle, 2001). Indeed, our own in vitro analysis of two such mutants [PrP(H187R) and PrP(E200K)] by both protease-protection and biotinylation assays confirmed this conclusion (Supplemental Figure S4). How such mutations lead to disease is unknown. Given that $\text{CtmPrP}$ is capable of causing disease, we decided to ask whether in vivo, any of the non-HD mutants might somehow affect $\text{CtmPrP}$ metabolism.

An initial screen using the biotinylation assay of most of the inherited disease-causing PrP mutants suggested that several (but not all) mutants have somewhat increased biotinylated products (unpublished observations). Among these, PrP(H187R) was especially obvious, and we therefore focused further efforts on this mutant. Our earlier analysis of PrP(H187R) had shown that relative to wild-type PrP, a greater proportion of the mutant is detergent insoluble, and this population is specifically enriched in immature glycosylated, EndoH-sensitive, intracellularly localized species (Ashok and Hegde, 2009). Because these properties are in many ways similar to the behavior of $\text{CtmPrP}$ (Figure 5), we asked if the mutant-enriched detergent insoluble fraction contained $\text{CtmPrP}$ as judged using our biotinylation assay. When PrP$\text{Bio}$ and PrP(H187R)$\text{Bio}$ lysates were fractionated on the basis of detergent solubility and probed for biotinylated species, nearly all of the biotinylated species in both samples was insoluble, and clearly increased for PrP(H187R)$\text{Bio}$ (Figure 7B). This difference was consistently seen regardless of how it was visualized: PrP immunoprecipitates probed with Avidin-HRP, and Avidin-pulldowns probed with anti-PrP both showed increased biotinylated products for PrP(H187R)$\text{Bio}$ (Figure 7B). This suggested that the total amount of $\text{CtmPrP}$ (and cyPrP) was greater for PrP(H187R) than for PrP. Yet, in vitro translocation analysis of PrP(H187R) did not show increased production of either $\text{CtmPrP}$ or cyPrP (Supplemental Figure S4). To resolve this discrepancy, we pulse-labeled cells and used Avidin-pull-downs to visualize the cyPrP and $\text{CtmPrP}$ being produced by PrP$\text{Bio}$ and PrP(H187R)$\text{Bio}$. An autoradiograph of the pulldowns showed that both the wild type and mutant generate comparable amounts of biotinylated species after a 30 min pulse labeling (Figure 7C). When the exact same samples were instead blotted with anti-PrP antibodies to visualize total cellular levels of biotinylated product, PrP(H187R)$\text{Bio}$ clearly bud ~2–3-fold more than PrP$\text{Bio}$. Thus, PrP and PrP(H187R) produce biotinylated species (i.e. $\text{CtmPrP}$ and cyPrP) at similar rates, consistent with in vitro translocation analyses; yet, the mutant has higher steady state levels. This strongly suggested that the turnover of the mutant $\text{CtmPrP}$ and cyPrP is slower compared with wild-type $\text{CtmPrP}$ and cyPrP.

To test this directly, we performed pulse-chase analysis (Figure 7D). When total PrP was analyzed by anti-PrP immunoprecipitation, a subtle difference was observed between PrP$\text{Bio}$ and PrP(H187R)$\text{Bio}$ wherein some bands were slightly overrepresented in the chase sample of the mutant (as has been previously observed; Ashok and Hegde, 2009). Pulldowns with Avidin of these same samples revealed the overrepresented species to be biotinylated. While the biotinylated species were reduced by ~50% at the 7 h chase time for PrP$\text{Bio}$, it was hardly diminished (~20%) for PrP(H187R)$\text{Bio}$. Although the absolute amount of biotinylated species varied somewhat from experiment to experiment, their selective stabilization with the mutant was consistently observed (Supplemental Figure S5). Thus, we conclude that the H187R mutation has the effect of stabilizing the turnover of $\text{CtmPrP}$, and possibly also cyPrP.

To further validate this conclusion, we asked whether improving signal sequence efficiency, which reduces $\text{CtmPrP}$ generation, would at least partially normalize the steady-state levels of detergent-insoluble immature species of PrP(H187R). Indeed, replacement of the PrP signal sequence with that from Pti did reduce the detergent insoluble, immature glycosylated species of PrP(H187R), with a corresponding increase in mature glycosylated soluble forms (Figure 7E). Had these mutant-specific aberrant species been generated from fully translocated forms of PrP (e.g., via their misfolding), it is difficult to see how they could have been partially rescued by improving signal sequence efficiency.

Thus, we conclude that among the aberrant species generated by PrP(H187R), $\text{CtmPrP}$ is likely to be a contributor, despite the fact that the mutation does not increase $\text{CtmPrP}$ production. Rather, it appears that the mutation selectively stabilizes, at least partially, the $\text{CtmPrP}$ form to increase its steady state levels and perhaps its trafficking. Given the association between $\text{CtmPrP}$ and neurodegeneration, it is attractive to speculate that at least one mechanism contributing to disease caused by these mutants is via increased levels of $\text{CtmPrP}$, which among other things may titrate away key cellular factors important for neuronal function (Chakrabarti and Hegde, 2009).

Although we have not yet done a complete analysis of all mutants or many cell types, we are able to confirm that the selective $\text{CtmPrP}$ stabilization is seen in other cell types and for at least one other disease-causing mutation (Supplemental Figure S6). Hence, in HEK293, we again observed increased biotinylated glycosylated PrP(H187R)$\text{Bio}$. In this case, the unglycosylated species was not noticeably increased, perhaps because the cytosolic quality control systems are somewhat different in these cells. In addition to H187R, another disease-causing mutant, PrP(E200K), also showed selective increase in glycosylated biotinylated species (Supplemental Figure S6). Similar results were obtained in HeLa and HT1080 cells (data not shown). Thus, mutants in the PrP C terminus can selectively influence the metabolism of $\text{CtmPrP}$, resulting in its increased levels at steady state. Importantly, not all mutations in the C terminus can be explained by this mechanism. For example, we found that PrP(V210I) is unchanged in the amount of $\text{CtmPrP}$ at steady state (data not shown), despite the fact that relative to wild type, a substantial amount of it is immature and detergent insoluble (Ashok and Hegde, 2009). Thus, there appear to be multiple effects of mutations in the C-terminal domain, and the relatively crude parameters of detergent insolubility and immature glycans cannot be used to equate all of the mutants. However, the more precise compartment-restricted biotinylation assay for $\text{CtmPrP}$ allowed us to reveal a novel mechanism by which at least some mutants of PrP influence its metabolism.

Conclusions

We have illustrated the ability of site-specific biotinylation to covalently mark the cytosolically exposed population of a protein ordinarily directed to the secretory pathway. In the case of PrP, this method could be combined with an ER-specific modification (glycosylation) to selectively identify and track the low-abundance but biologically
important CtmPrP species. This method substantially improves upon other methods to detect CtmPrP, which are either biochemically cumbersome (e.g., topology assays on ER microsomes) or based on a faulty premise (such as lack of signal sequence processing). The method is bio-orthogonal, has a high signal-to-noise, can be readily applied to other systems, and works in formats that allow both steady state analysis and pulse-chase studies. While there are endogenous biotinylated proteins, they do not seem to pose a major source of background in our biochemical assays. Furthermore, it is straightforward to pre-enrich for one’s protein of interest (e.g., with immunoprecipitation or ConA in the case of CtmPrP) before detecting biotinylation to markedly improve signal-to-noise. While direct spatial visualization by fluorescence (e.g., using fluorophore-labeled Avidin on fixed cells) is also feasible, this remains to be investigated for the PrP system.

The utility of this method is exemplified by several new insights into CtmPrP biosynthesis and metabolism. Not only were we able to directly detect the CtmPrP population of wild-type PrP, but we also could show that it is a relatively stable membrane protein that samples much of the secretory pathway including the cell surface. The finding that CtmPrP is not rapidly degraded from the ER is especially noteworthy, because it suggests that this species may not be recognized as aberrant by the cellular quality control machinery. Indeed, the elements involved in CtmPrP production (the signal sequence and central hydrophobic domain) are highly conserved, and it is possible that it has a normal function that remains to be elucidated. In this scenario, only under conditions of excess production or stabilization in certain cell types would CtmPrP be detrimental. The ability to set up a system for selectively isolating this population (e.g., by ConA followed by Avidin affinity steps) now paves the way to determining whether its levels vary in vivo in a cell type-specific, aging-dependent, or disease-dependent manner in mice engineered to express PrPΔ530 and BirA. That CtmPrP is degraded in lysosomes may suggest a means by which accumulation of transmissible prions could lead to increased CtmPrP (Hegde et al., 1999). In this view, prion accumulation would lead to lysosomal dysfunction, which would delay CtmPrP degradation. Being able to follow CtmPrP metabolism should permit this and related hypotheses to be tested.

At least in cell culture, the ability to readily detect CtmPrP revealed an unexpected mechanism leading to its increased levels. Here, mutants in the C-terminal domain of PrP act preferentially on the CtmPrP form to stabilize it. This finding would have been very difficult to obtain by other means given the rather subtle effect and the difficult biochemical properties of CtmPrP. Whether this stabilization is due to increased aggregation, altered trafficking, or another reason remains to be determined. However, the increase, while relatively small, is nonetheless likely to be important given that even small increases in CtmPrP seem to be capable of causing neurodegeneration in transgenic mouse models (Hegde et al., 1998; Hegde et al., 1999; Rane et al., 2010).

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