Maturation of the trans-Golgi Network Protease Furin: Compartmentalization of Propeptide Removal, Substrate Cleavage, and COOH-terminal Truncation

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Abstract. We have cloned a bovine cDNA encoding the trans-Golgi network (TGN) protease furin and expressed it via recombinant vaccinia viruses to investigate intracellular maturation. Pulse-chase labeling reveals that the 104-kD pro-furin bearing high mannose N-glycans is rapidly processed into the 98-kD protease whose N-glycans remain sensitive to endoglycosidase H for a certain period of time. Furthermore, in the presence of brefeldin A, pro-furin cleavage occurs. From these data we conclude that the ER is the compartment of propeptide removal.

Studies employing the ionophore A23187 and DTT show that autocatalysis is Ca\(^{2+}\) dependent and that it does not occur under reducing conditions. Pro-furin produced under these conditions never gains endo H resistance indicating that it is retained in the ER. Coexpression of furin with the fowl plague virus hemagglutinin in the presence of brefeldin A and monensin reveals that furin has to enter the Golgi region to gain substrate cleaving activity.

N-glycans of furin are sialylated proving its transit through the trans-Golgi network. A truncated form of furin is found in supernatants of cells. Truncation is inhibited in the absence of Ca\(^{2+}\) ions and in the presence of acidotropic agents indicating that it takes place in an acidic compartment of cells.

Comparative analysis with furin expressed from cDNA reveals that the truncated form prevails in preparations of biologically active, endogenous furin obtained from MDBK cells. This observation supports the concept that secretion of truncated furin is a physiological event that may have important implications for the processing of extracellular substrates.

Many bioactive peptides and proteins like several peptide hormones, growth factors, coagulation factors, or receptors on the plasma membrane are synthesized as precursors which have to be cleaved by cellular endoproteases to gain their full biological potential (Douglass et al., 1984; Sossin et al., 1987; Furie and Furie, 1988; Yoshimasa et al., 1990; Hosaka et al., 1991). Although they differ with respect to their cell type-specific expression and their mode of secretion, they share some common features. Activation cleavage takes place at the carboxyterminal arginine of a di-, tri-, or tetrabasic recognition peptide and this process is mediated by Ca\(^{2+}\)-dependent endoproteases in a late compartment of the exocytic pathway of cells (Docherty and Steiner, 1982; Barr, 1991). This concept has been developed early in the evolution of eukaryotes because even the unicellular organism Saccharomyces cerevisiae secretes bioactive peptides like the alpha mating factor or the killer toxin by cleavage of their protein precursors at Lys-Arg or Arg-Arg sites (Bostian et al., 1984; Julius et al., 1984a). While maturation and activation of these proproteins have been studied in detail, information on the nature and characteristics of the activating enzymes was rare.

An important step in the identification of the proteases was the discovery of the yeast endoprotease kexin which mediates the cleavage of pro alpha-factor and pro killer toxin (Julius et al., 1984a; Fuller et al., 1988, 1989). Homologues of kexin in higher eukaryotes could then be found because all of these enzymes share a subtilisin-like catalytic domain. Furin, encoded by the fowl plague virus gene, the fes/fps upstream region gene (also called PACE for paired basic amino acids cleaving enzyme), first of the mammalian subtilisins to be discovered (Roebroeck et al., 1986; Van den Ouweland et al., 1990; Smeekens and Steiner, 1990; Creemers et al., 1992)

Abbreviations used in this paper: BFA, brefeldin A; FPV, fowl plague virus; HA, hemagglutinin; MDBK, Madin Darby Bovine Kidney; NA, neuraminidase; PC, prohormone convertase; TGN, trans-Golgi network.
are confined to endocrine and neuroendocrine tissues (Seidah et al., 1990; Smeekens et al., 1991) where they cleave prohormones like proinsulin or proopiomenocortin (Benjannet et al., 1991; Thomas et al., 1991); PC4 mRNA has only been detected in testis (Nakayama et al., 1992), and PC6 (also called PC5) expression is mainly found in intestinal cells (Nakagawa et al., 1993). Besides the subtilisin-like catalytic domain, these enzymes share an NH2-terminal signal sequence and a prodomain which for some of these proteases has been shown to be cleaved as it is the case with subtilisin (Power et al., 1986) and kexin (Brenner and Fuller, 1992). On the other hand, there are important differences between the eukaryotic subtilisins. While most of them lack a hydrophobic anchor domain, furin and kexin are integral membrane proteins. Furthermore, the prohormone convertases PC1/3 and PC2 are sorted into and act in the regulated secretory pathway of endocrine cells, whereas furin acts in the constitutive secretory pathway present in all cells.

According to the intracellular targeting and tissue distribution furin and the prohormone convertases have different substrates and cleavage site specificities: substrates of furin are activated at the recognition motifs Arg-X-Lys/Arg-Arg or Arg-X-X-Arg (Molloy et al., 1992) in contrast to PC1/3, PC2, and kexin cleaving at Lys-Arg- or Arg-Arg-sites (Fuller et al., 1988; Thomas et al., 1991). Endogenous furin substrates can either be soluble proteins like pro-nerve growth factor (Bresnahan et al., 1990), pro-von Willebrand factor (Van de Ven et al., 1990; Wise et al., 1990) or membrane bound glycoproteins like pro-insulin receptor (Robertson et al., 1993) which are transported via the constitutive secretory pathway. Moreover furin plays an important role in the pathogenicity of certain bacteria and viruses. Pseudomonas exotoxin A and Bacillus anthracis protective antigen can be activated by furin although the cellular compartment of cleavage of these exogenous substrates has not been defined yet (Molloy et al., 1992; Moehring et al., 1993). Viruses, on the other hand are intracellular pathogens using the cellular machinery for production of their progenies. Some of the enveloped viruses, for example paramyxoviruses like the pathogenic Newcastle disease viruses (Gotoh et al., 1992; Moehring et al., 1993) and human parainfluenza virus type III (Ortmann et al., 1994), human immunodeficiency virus (HIV) (Hallenberger et al., 1992), or fowl plague virus (FPV) (Stieneke-Gröber et al., 1992) bear integral glycoproteins that are activated by furin before virions leave their host cells in a budding process (for review see Klenk et al., 1992). For the hemagglutinin (HA) of FPV it has been shown that the endogenous cleaving activity of the permissive Madin Darby Bovine Kidney (MDBK) cell line has the same properties as furin, and in enriched Golgi fractions furin could be detected by immunoblotting (Stieneke-Gröber et al., 1992).

Furin is synthesized as a prepro-enzyme at the rough endoplasmic reticulum and later on it is concentrated in the trans-Golgi network (TGN) (Bresnahan et al., 1990; Misumi et al., 1991; Molloy et al., 1994; Schäfer, W., M.-L. Kruse, M. Vey, S. Bergbøker, J. Seiler, H. F. Kern, H.-D. Klenk, and W. Garten, manuscript submitted for publication). Therefore, maturation and transport processes that happen in between are of special interest, but knowledge on temporal and spatial relationships between these stages of furin maturation is still incomplete. Previous studies have shown that proteolytic removal is an autocatalytic process (Leduc et al., 1992) but different opinions on the cellular compartment in which this reaction takes place, based on soluble furin mutants with attached KDEL sequences, exist (Rehemtulla et al., 1992; Molloy et al., 1994). Pulse-chase studies favor the ER as the compartment of propeptide removal (Molloy et al., 1994). After activation, wild-type furin accumulates in the TGN (Molloy et al., 1994; Schäfer, W., M.-L. Kruse, M. Vey, S. Bergbøker, J. Seiler, H. F. Kern, H.-D. Klenk, and W. Garten, manuscript submitted for publication), while truncated furin mutants are secreted from cells (Hatsuzawa et al., 1992; Molloy et al., 1992). Finally, the fate of furin has not been clarified yet, but there are indications that the membrane-bound enzyme is converted into a soluble form that can be found in supernatants of overexpressing cells (Rehemtulla et al., 1992; Vidricaire et al., 1993; Molloy et al., 1994). It should be emphasized that all of our knowledge on furin biosynthesis has been derived from studies on the overexpressed protease.

We report here on the spatial and temporal relationship between synthesis, autocatalytic propeptide removal, transport, substrate cleavage, and secretion of furin. Our studies have been carried out on bovine and human furin overexpressed from cDNA and on endogenous bovine furin enriched from Golgi fractions of MDBK cells.

**Materials and Methods**

**Cell Culture**

African green monkey kidney (CV-1) cells and MDBK cells were grown in Dulbecco's medium supplemented with 10% fetal calf serum. For production of recombinant vaccinia viruses and all labeling procedures, cells were grown on 60-mm-diam culture dishes (GIBCO, Eggenstein, Germany). Human TK-143 cells were grown in Dulbecco's medium supplemented with 5% fetal calf serum and 25 μg/ml of 5-bromo-2'-deoxyuridine (Sigma Chem. Co., Deisenhofen, Germany). The Western Reserve strain of vaccinia virus was propagated in CV-1 cells.

**Isolation of Endogenous MDBK Furin and Detection by Immunoblotting**

Golgi fractions were prepared from MDBK cells and furin enriched by chromatography on FPLC as described previously (Stieneke-Gröber et al., 1992). For immunodetection an aliquot of the Superose fractions and lysates of CV-1 cells harvested 5 h after infection with recombinant vaccinia virus VV:hfur (multiplicity of infection [m.o.i.] = 10 pfu/cell) were separated on 12% gels under reducing conditions. Proteins were transferred to nitrocellulose and detected with a furin specific antiserum from rabbits.

**Amplification of a Human Fur Gene Fragment**

DNA of a recombinant vaccinia virus encoding the human furin cDNA (Bresnahan et al., 1990) was isolated by standard procedures (Macket et al., 1985) and used as template for PCR amplification of a part of the region coding for the subtilisin-like domain. The following primers were used: 5'-CACGGCACACGGTGTC~AAGT-3' corresponding to nucleotides 580 to 605 and 5'-CAGGTGTTGCATGTCCCGCCATGTGA-3' complementary to nucleotides 1163 to 1188 of the coding region of the human furin. For amplification a commercially available kit (Perkin-Elmer Cetus Instruments, Weiterstadt, Germany) was used and reactions were performed in a thermal cycler (Perkin-Elmer Cetus Instruments) for 30 cycles of denaturation (94°C, 2 min), annealing (55°C, 2 min), and extension (72°C, 1.5 min). After electrophoresis the DNA fragment was extracted from the agarose gel and used as template for production of digoxigenin tagged probes employing a random primed DNA labeling kit (Boehringer, Mannheim, Germany).
Cloning of Bovine Fur cDNA from an MDBK cDNA Library

About 2 × 10^6 clones of a ZAP-Express cDNA library of MDBK cells (Stratagene, Heidelberg, Germany) were screened with the human fur-1 probe using standard protocols for blotting and hybridization techniques. Bovine fur-positive clones were visualized by a nonradioactive detection kit employing chemiluminescent reagents (Boehringer). Five fur-recombinant clones were selected and plaque purified and their cDNA sequences determined by standard protocols for blotting and hybridization techniques. One of these clones contained a full-length fur cDNA from which the 2.8-kb Smal fragment bearing the furin open reading frame was subcloned into a pBluescript vector (Fig. 1 A, lane V'E'hfur) for production of recombinant vaccinia viruses. The DNA sequence of both strands of the Smal fragment was determined by the dye-deoxyribonucleotide chain termination sequencing method (Sanger et al., 1977).

Construction of the Bovine Furin Recombinant Vaccinia Virus

Subconfluent CV-1 cells were infected with the Western Reserve strain of vaccinia virus (moi. 0.05 pfu/cell) 2 h before cells were lipofected with the pSc1/bfur construct. 36 h after infection virus stocks were prepared, and recombinant viruses were selected on TK- cells in the presence of 25 μg 5-bromodeoxyuridine (Sigma Chem. Co.) per ml. After three rounds of plaque purification on CV-1 cells, virus stocks were prepared as previously described (Macket et al., 1984; Chakrabarti et al., 1985).

Metabolic Labeling of Infected Cells, Immunoprecipitation, Carbohydrate Analysis, and SDS-PAGE of Precipitated Proteins

In all single expression experiments, CV-1 or MDBK cells were infected with wild-type vaccinia virus, bovine furin, or human furin recombinant vaccinia viruses at a moi. of 10 pfu/cell. For coexpression, CV-1 cells were infected with bovine furin recombinant vaccinia virus at a moi. of 5 pfu/cell and either FPV neuraminidase or FPV hemagglutinin recombinant viruses at a moi. of 5 pfu/cell. At 3.5 h after infection cells were starved for methionine and either FPV neuraminidase-negative or FPV neuraminidase-free Dulbecco's modified Eagle's medium for 1 h. When the ionophore A23187 was applied, methionine-free Dulbecco's minimal essential medium lacking Ca^2+ and Mg^2+ was used. Cells were pulse labeled with 100 μCi of L-[35S]-methionine (Amersham-Buchler, Braunschweig, Germany) per ml medium (1,000 Ci/mmol) for different periods of time and then either lysed immediately or after incubation in the respective media containing unlabeled methionine (2 mM). 1.9% was done in 1 ml of RIPA buffer (1% Triton X-100, 1% sodium deoxycholate, 0.1% SDS, 0.15 M NaCl, 20 mM Tris, pH 7.5, 10 mM EDTA, 1 mM phenylmethylsulfonyl fluoride [Sigma Chem. Co.], 0.5 μg/ml aprotinin, and 10 mM 2-iodoacetamide [Sigma Chem. Co.]). Before immunoprecipitation, nuclei were removed by centrifugation at 10,000 g for 30 min at 4°C. For detection of secreted furin cell supernatants were cleared from cell debris and diluted in a 1:1 PA buffer (0.1 M NaCl, 50 mM Tris, pH 7.5, 1% NP-40, 1 mM EDTA, 20 μM phenylmethylsulfonyl fluoride, 1% Triton X-100). For immunoprecipitation, cleared lysates and supernatants of single expression experiments were incubated with 2 μl of furin specific antiserum from rabbits and 40 μl of protein A-Septarose suspended 1/1 in RIPA buffer. Lysates of coexpression experiments were divided into two halves, one of which was incubated with the furin specific antiserum and the other one with a FPV specific antiserum. For analysis of carbohydrate side chains of the immunoprecipitated proteins, endoglycosidase H and N-glycanase F digestions were carried out as previously described (Roberts et al., 1993). Proteins were then subjected to SDS-PAGE on 10% gels under reducing conditions and visualized by fluorography.

Application of Modulating Compounds

For transport inhibition studies, infected cells were incubated either with brefeldin A (Sigma Chem. Co.) or monensin (Sigma Chem. Co.) at a concentration of 2 μg/ml during starvation, pulse, and chase periods. To study the effect of different Ca^2+ concentrations on furin activity, procedures were carried out in medium supplemented with ionophore A23187 (Boehringer) at a final concentration of 0.25 μM. Dithiothreitol (DTT) was added to media at 1 mM to inhibit disulfide bridge formation during starvation and labeling periods. To raise the pH of acidic cellular organelles, cells were either treated with ammonium chloride or chloroquine (Sigma Chem. Co.) at final concentrations of 30 mM and 80 μM, respectively. To test the effect of Ca^2+ removal on truncation and secretion of furin, the ionophore was added only during the chase period.

Results

Comparison of Bovine and Human Furin

We have isolated the endogenous furin protease from Golgi fractions of MDBK cells as described previously (Stieneke-Gröber et al., 1992) and compared it with overexpressed human and bovine furin (Fig. 1). The 85-kD protein band of the endogenous furin (Fig. 1 A, lane MDBK) differs significantly from the 94- and 100-kD doublet of the human protease (Fig. 1 A, V'E'hfur). To find out if this difference is caused by mutations in the coding region of the bovine fur gene we have cloned its cDNA from an MDBK library and determined the nucleotide sequence (EMBO Data Bank accession number X75956). The deduced amino acid sequence revealed homology with human furin except for a proline-rich region upstream of the membrane anchor comprising amino acids 686 and 696 of human furin (Table I). In bovine furin this region is extended by three additional amino acid residues, and two of these are prolines. MDBK cells were infected to study the expression of furin (Fig. 1 B). Cells were either infected with human (lanes V'E'hfur) or bovine furin recombinant viruses (lanes Vhfur), and proteins were pulse labeled for 45 min (Fig. 1 B, lanes C) or labeled for 45 min and then chased for 4 h with cold methionine (Fig. 1 B, lanes S). Furin proteins were precipitated with pVhfur antiserum and resolved by SDS-PAGE on 10% gels.

Figure 1. Endogenous bovine furin is smaller than overexpressed, membrane-bound forms of human and bovine furin but similar to their secreted forms. (A) Immunodetection of endogenous bovine and overexpressed human furin. Golgi fractions of MDBK cells were enriched and, after solubilization, subjected to FPLC on a Superose 12 column as previously described (Stieneke-Gröber et al., 1992). An aliquot of a furin containing fraction was prepared for SDS-PAGE by boiling in sample buffer containing β-mercaptoethanol (lane MDBK). For comparison with overexpressed human furin CV-1 cells infected with recombinant vaccinia viruses VV; hfur (moi. = 10) were lysed in sample buffer (lane V'E'hfur) 5 h p.i. Proteins were separated on a 12% gel and blotted onto nitrocellulose. Furin proteins were detected with a furin specific antiserum from rabbits. (B) Fluorogram of radiolabeled bovine and human furin immunoprecipitated from cell lysates and supernatants of overexpressing cells. MDBK cells were infected with recombinant vaccinia virus encoding human (Vhfur) or bovine (V'E'hfur) furin (moi. = 10). At 3.5 h after infection cells were either pulse labeled for 45 min (lanes C) with 35S-methionine or labeled for 45 min and chased for 4 h with cold methionine (lanes S). Furin was then immunoprecipitated from pulse labeled cell lysates (lanes C) and from supernatants of the pulse-chase-labeled cells with a furin specific antiserum and resolved by SDS-PAGE on 10% gels. The arrows indicate positions of the molecular mass.
ties migrating at 100 and 94 kD. Previous studies on human
upper band had a relative molecular mass of 104 kD and the
very similar. Bovine furin appeared as a doublet of which the
pro-furin, that is converted into the mature furin protease
furin (Bresnahan et al., 1990; Van de Ven et al., 1990; Wise
migration of bovine furin. Again bovine furin showed reactivity
with this antiserum proving that human and bovine furin are
very similar. Bovine furin appeared as a doublet of which the
upper band had a relative molecular mass of 104 kD and the
lower band migrated at 98 kD, whereas the doublet bands ob-
erved with human furin had higher electrophoretic mobili-
ties migrating at 100 and 94 kD. Previous studies on human
furin (Bresnahan et al., 1990; Van de Ven et al., 1990; Wise
et al., 1990; Leduc et al., 1992; Creemers et al., 1993) had
shown that both of these proteins are encoded by the fur open
reading frame: the larger protein represents the proenzyme,
pro-furin, that is converted into the mature furin protease
(Fig. 1 A, lanes C, lower band) by autoproteolytic removal of
the propeptide. The differences in electrophoretic mobili-
ties of human and bovine furin were not cell type dependent
as the expression in CV-1, NRK, and MDBK cells indicated
data not shown). After removal of N-linked carbohydrate side chains by digestion with N-glycanase F (data not
shown), the differences were still detectable. Thus, there
may be conformational differences or other posttranslational modifications which account for the lower electrophoretic
mobility of bovine furin.

Although the membrane bound, intracellular form of the
overexpressed bovine furin has an even higher molecular
mass than its human counterpart and the data of its nucleo-
tide and deduced amino acid sequence showed no relevant
mutation accounting for the observed difference between the endogenous and the overexpressed protease there is an im-
portant link between these data which is obvious after im-
munoprecipitations of the truncated and secreted furin pro-
teins. Truncated human and bovine furin seem to be clamped off their membrane anchors at a specific site in the luminal domain because they have similar molecular masses (Fig. 1
A, lanes S) except for the small difference that could be de-
tected for the membrane bound proteins (Fig. 1 B, lanes C)
before. Truncation releases an 85-90-kD protein which can be
found in the supernatants of cells (Fig. 1 B, lanes S). Thus,
the truncated furin in supernatants of overexpressing cells seems to have the same molecular mass as the endoge-
nous furin of MDBK cells.

In summary, the cloned and overexpressed bovine furin,
when expressed in mammalian cells, seems to be autoca-
tactically active and it has a higher molecular mass than hu-
man furin. In addition, these results suggest that the endoge-
nous furin from MDBK cells was isolated in its truncated
form.

Table I. Differences in the Amino Acid Sequences of Bovine and Human Furin

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Single letter code has been used for abbreviation of amino acid residues. Amino acid sequence of human furin was taken from Van den Ouweland et al. (1990).

Propeptide Removal Occurs Rapidly
and Precedes Transport to Golgi Apparatus
and COOH-Terminal Truncation

Little is known about the temporal and spatial relationship
of maturation and transport of furin. Earlier reports postu-
lated a post ER compartment for autoprocessing and sub-
strate cleavage (Rehemtulla et al., 1992). Studies on rat
furin, on the other hand, showed that the N-linked carbo-
hydrates of pro-furin and cleaved furin were endo H sensitive
and were converted only after several hours into complex
glycans (Misumi et al., 1991). Additionally, contradictory
conclusions were drawn from studies on soluble furin mu-
ant with attached KDEL sequences (Rehemtulla et al.,
1992; Molloy et al., 1994). Thus, the compartment of auto-
processing has not been clearly defined.

Pulse-chase labeling studies have been performed before
on epitope-tagged furin (Molloy et al., 1994) and wild-type
human furin (Creemers et al., 1993), and we have used a
similar approach to study kinetics of propeptide removal of
native bovine and human furin. In addition, we analyzed the
status of the N-linked glycans of furin during the maturation
processes to link the kinetics of propeptide removal with data
on transport through the compartments of the secretory path-
way (Fig. 2 A). Only the data observed with bovine furin are
shown, but similar results were obtained on human furin.
Autocatalysis is a posttranslational event as indicated by the
observation that most of the immunoprecipitated furin was
labeled in its pro-form during a 10-min pulse (Fig. 2 A, panel
0'). In addition to the proenzyme, cleaved furin could be de-
tected indicating that propeptide removal had already begun.
Both of the precipitated furin bands shifted after treatment
with endoglycosidase H to the molecular mass of the
deglycosylated forms which had been generated by digestion
with N-glycanase F. This means that all N-linked sugars were
of the high mannose type and, thus, autoprocessing probably
took place in the ER. During the following chase periods
(Fig. 2 A, 10', 20', and 30') pro-furin was completely
converted into cleaved furin. The half time of the autocatalytic
reaction could not be measured precisely, but between 0 and
Figure 2. Autoproteolytic cleavage precedes intracellular transport and secretion of furin. Fluorograms of bovine furin: CV-1 cells were infected with VV:bfur and metabolically labeled with 35S-methionine 4.5 h after infection. After labeling furin proteins were immunoprecipitated from lysates with a furin-specific antiserum and analyzed by SDS-PAGE on 10% gels and fluorography. (A) Autoproteolytic cleavage of pro-furin and intracellular transport of cleaved furin. Pulse-chase labeling: cells were labeled for 10 min and either lysed immediately (0') or incubated in chase medium for the indicated periods (10, 20, 30, 40, and 60 min). Aliquots of the proteins were digested with endoglycosidase H or N-glycanase F before SDS-PAGE to study the structure of carbohydrate side chains. (B) Secretion of mature furin. Pulse-chase labeling: cells were labeled for 45 min and either lysed immediately (0 h) or incubated in chase medium for the indicated periods (1, 2, 3, or 4 h). Cell lysates as well as supernatants were incubated with furin-antiserum to precipitate the intracellular furin and the secreted, truncated form of mature furin. (C) Truncation of mature furin takes place inside the cells. Cells were labeled for 2 h and chased for 1 h. Furin was immunoprecipitated from cell lysates and aliquots were digested with endoglycosidase H or N-glycanase F. Arrow indicates position of molecular mass standard; n, position of pro-furin; ~//>, autocatalytically cleaved furin with endo H-sensitive carbohydrate side chains; o, cleaved furin after truncation of the COOH-terminal, membrane-spanning region; d, deglycosylated form of the marked protein.

10 min of chase half of the labeled pro-furin was converted into the cleaved form. Endo H treatment clearly showed that furin still had high mannose glycans indicating that after propeptide removal the cleaved form only slowly left the ER. After 40 and 60 min of chase, again a high molecular mass furin band appeared, which did not shift after endo H treatment demonstrating that N-linked carbohydrates of this protein had been converted into complex types. This larger band now represented cleaved furin because it shifted to the position of the deglycosylated, cleaved furin after digestion with N-glycanase F. These results show that autoprocessing of pro-furin is an early event starting a few minutes after synthesis. Only the cleaved form slowly reaches the medial and late Golgi compartment, where its N-linked glycans are converted into complex types.

Fig. 2 B monitors furin maturation during longer chase periods. In this experiment supernatants of furin expressing cells were also incubated with the anti-furin antibody to check if a soluble form of furin had been secreted into the medium as has been reported in earlier studies (Wise et al., 1990; Rehemtulla et al., 1992). After a 45-min pulse without chase, the cleaved furin band dominated over pro-furin, because the labeling period had exceeded the half time of autocatalysis about three times (Fig. 2 B, bottom). After 1 h of chase, pro-furin totally disappeared, and the doublet now consisted of high and low molecular mass cleaved furin. The endo H-resistant furin did not run as a distinct band, but a smear of labeled proteins could be detected between 98 and 110 kD hinting at different amounts of sialic acid attached to its oligosaccharides. In addition to these two forms, a smaller band with an estimated molecular mass of 90 kD could be precipitated from the cell lysates and a protein of similar size could also be detected in the supernatants of the cells (Fig. 2 B, top). During the following chase periods, the intracellular level of labeled furin decreased markedly and after 4 h it could no longer be detected. In contrast, the
amount of the soluble furin protein in the supernatant increased to the same extent which proved that furin was efficiently secreted from the cells.

The three forms of furin detected after propeptide removal could be precipitated from the cell lysates (Fig. 2 C) after 2 h labeling and 1-h chase period. The fluorogram showed two diffuse bands extending from 98 to 110 kD and from 80 to 90 kD and one sharp band at 98 kD. Endo H treatment revealed that the two heterogeneously migrating forms of furin had complex carbohydrate side chains, whereas the distinct band shifted from 98 kD in the untreated sample to ~92 kD. After deglycosylation, the larger band again shifted to the position of the 92-kD deglycosylated cleaved furin (see Fig. 2 A). The deglycosylated form of soluble furin, on the other hand, had an estimated molecular mass of 76 kD. From these results we conclude that the truncated, secreted furin was produced from the mature, membrane bound form by proteolytic cleavage and that this process might have taken place inside the cell in a late compartment of the exocytic pathway. These results indicate that the 85-kD furin isolated from MDBK cells represents the COOH-terminally truncated form of bovine furin.

In conclusion, pulse-chase labeling experiments revealed that propeptide removal from the 104-kD precursor had complex carbohydrate side chains starts soon after synthesis. After 40-min chase, pro-furin completely disappears. Cleaved furin bearing endo H-sensitive N-glycans with a molecular mass of 98 kD is then transported to the Golgi region, and about 50 min after pulse complex carbohydrate side chains can be found on the protein. The endo H-resistant, cleaved furin now migrating at 104 kD is then COOH-terminally truncated and, 4 h after the pulse, all of the labeled furin is found in the supernatant of cells as a 80-90-kD protein. This maturation pathway of furin is schematically shown in Fig. 8. It is now evident that maturation, transport, and secretion of bovine furin are performed in individual, consecutive steps which seem to be precisely scheduled and may be confined to distinct compartments in the exocytic pathway of cells.

**Mature Furin Is Sialylated**

To find out if the increase in the apparent molecular mass of furin from 98 kD to ~104 kD (Fig. 2 A and B) was due to sialylation, furin was coexpressed with the neuraminidase (NA) of FPV (Fig. 3). FPV NA hydrolyzes the α2-3 glycosidic bond between terminal sialic acid and galactose on glycoproteins and glycolipids. It is a membrane bound glycoprotein, matures in the exocytic pathway of cells and is transported to the plasma membrane. After coexpression with NA, furin had a molecular mass of 98 kD and was not sensitive to endo H treatment (Fig. 3, lanes 4–6). Thus, the increase in molecular mass observed after single expression results from the attachment of neuraminic acid to the carbohydrate side chains. After N-glycanase F treatment, furin obtained by single expression had the same electrophoretic mobility as furin obtained by coexpression with NA (Fig. 3, compare lane 3 with lane 6) indicating that sialic acid is attached to N-linked glycans but not to O-linked glycans. The observations that furin is sialylated indicates that furin enters the TGN where sialyltransferases are localized.

![Figure 3. Mature furin is sialylated. Fluorogram of bovine furin coexpressed with the fowl plague virus neuraminidase. 4.5 h after coinfection of CV-1 cells with vaccinia viruses encoding bovine furin (VV:bfur) and the neuraminidase of FPV (VVNA), cells were labeled with 35S-methionine for 1 h and chased for 1 h with cold methionine. Furin was immunoprecipitated from cell lysates and before SDS-PAGE on a 10% gel aliquots of proteins were digested with endoglycosidase H or N-glycanase F. Furin expressed in the absence of neuraminidase served as control. Arrow indicates position of the molecular weight marker; s, sialylated, autocatalytically cleaved furin; 0, autocatalytically cleaved furin lacking sialic acids; d, deglycosylated form of the marked protein.](http://jcb.rupress.org/content/jcb/127/6/1830.full.html)
with rabbit antiserum raised against FPV high marmose carbohydrate side chains; ~1, indicates autocatalytically cleaved furin with complex, sialylated sugar moieties; d symbolizes 0.1, or 1 mM CaCl₂. Cell lysates were divided and one half was incubated with antiserum specific for furin deglycosylated form of the marked protein.

Arrows on the left of fluorograms mark positions of molecular mass standards; arrows on the right mark positions of the precursor Furin labeled in the absence of Ca²⁺ ions (see A, lane 0) and digested with endoglycosidase H and N-glycanase F before SDS-PAGE. Arrows on the right mark positions of the precursor (HA) and cleavage products (HA₁ and HA₂) of FPV hemagglutinin; □, position of pro-furin; <, points to autocatalytically cleaved furin with high mannose carbohydrate side chains; 4, indicates autocatalytically cleaved furin with complex, sialylated sugar moieties; d symbolizes deglycosylated form of the marked protein.

These experiments underline the important role Ca²⁺ ions play not only in substrate processing but also in the autocatalytic cleavage reaction. In addition, pro-furin produced in the absence of Ca²⁺ is retained in an early compartment by a kind of quality control mechanism.

Inhibition of Pro-furin Processing Is Reversible
Retention of furin in the ER or early Golgi under conditions of Ca²⁺ depletion may result from irreversible aggregation as described for other misfolded proteins (Gething et al., 1986; Roberts et al., 1993). To check if inhibition of pro-furin autoprocessing is reversible, bovine furin was labeled in the absence of Ca²⁺ for 1 h and then chased for 1 h in the presence of Ca²⁺ ions (Fig. 5 A). No processing and no transport of pro-furin could be detected after pulse labeling (Fig. 5 A, left). After the chase, however, autocatalytic cleavage (Fig. 5 A, right) and transport of furin occurred. Thus, the transport block of furin can be released by the addition of Ca²⁺ ions.

Disulfide Bridge Formation Is Critical for Pro-furin Processing
Conformation of glycoproteins that enter the exocytic pathway of cells is mainly stabilized by disulfide bridges formed during and shortly after protein synthesis. We wanted to find out if disulfide bridge formation contributes to the correct processing of pro-furin, too. Studies on the folding of influenza hemagglutinin had employed the reducing agent DTT for reversible inhibition of oxidation of sulphydryl groups (Braakman et al., 1992). In analogy, we added the reducing agent to CV-1 cells expressing bovine furin (Fig. 5 B). In the presence of DTT, pro-furin showed no autoprocessing during a 1-h labeling period as could be demonstrated by the presence of the endo H-sensitive, 104-kD protein (Fig. 5 B, left). After extensive washing with DTT-free medium and an additional chase period in the absence of this chemical (Fig. 5 B, right) pro-furin was processed to the cleaved 98-kD furin. Compared to autoprocessing of nascent pro-furin kinetics of autocatalysis seemed to be slowed down after the release of the DTT shock (Fig. 1 A). This delay might reflect repair of incorrect disulfide bridges. The cysteine-rich region that precedes the transmembrane domain might play a critical role in these early folding processes. In conclusion, this experiment demonstrates that the oxidizing conditions in the ER are prerequisite for autoprocessing of pro-furin to occur.

Autocatalysis, but Not Substrate Cleavage, Takes Place in the Endoplasmic Reticulum
The results of pulse-chase labeling had shown that autoproteolytic processing probably occurred in the ER which is in contrast to previous studies postulating a post-ER compartment for cleavage as well as for autoprocessing (Rehemtulla et al., 1992). We therefore disintegrated the Golgi region by application of the fungal metabolite brefeldin A (BFA) (Lippincott-Schwartz et al., 1989), which, as a net result, blocked exit of the newly synthesized glycoproteins out of the ER (Klausner et al., 1992), and studied autoprocessing and substrate cleavage by coexpression with FPV hemagglutinin (Fig. 6 A). Pulse-chase labeling in the absence of BFA (Fig. 6 A, left) showed that during a 15-min pulse autocatalysis had already started but that hemagglutinin had not been cleaved by furin. There are at least three reasons why substrate cleavage had not occurred. Firstly, HA may not have acquired the correct conformation competent for cleavage by furin which may require trimerization or other
posttranslational events. Secondly, furin itself might not have the proper conformation. Pulse-chase experiments suggest that even after cleavage furin stays in the ER for some time, since it takes ~30 min to become endo H resistant (Fig. 2 A). Thirdly, some kind of chaperone may inhibit furin activity.

During the chase period, on the other hand, pro-furin disappeared and HA had been efficiently cleaved. Under these conditions of early vaccinia virus infection, not only the overexpressed bovine but also the endogenous furin of CV-1 cells is present, and we could not rule out that the endogenous furin activity contributed to the HA cleavage. Apart from that, this control experiment confirmed that HA cleavage under normal conditions takes place when HA has passed the Golgi apparatus. When coexpressed in the presence of BFA, on the other hand, newly synthesized furin and HA could not be exported from the ER, and posttranslational modifications normally performed in this compartment should have also taken place (Russ et al., 1991). Pulse (15

**Transport to the Golgi Apparatus Is Necessary for Furin Activity**

Early studies on the localization of furin demonstrated a concentration in the Golgi region (Bresnahan et al., 1990; Misumi et al., 1991) and recent findings show that it colocalizes with the trans-Golgi network protein TGN 38 (Molloy et al., 1994). The presence of sialic acid residues on the N-linked glycans of mature furin (Fig. 3) further underlines that a late compartment might be the site of accumulation and action of furin. Coexpression studies in the presence of monensin, which inhibits transport of proteins from cis- and medial-Golgi cisternae to the trans region should give an answer to the question in which compartment of the Golgi region furin can activate substrates. Pulse-chase analysis (Fig. 6 B) in the absence of monensin showed that, regardless of coexpression or single expression, furin and HA matured normally (Fig. 6 B, right top and bottom): furin was cleaved and extensively sialylated which was also true for HA. In the presence of monensin (Fig. 6 B, top left), pro-furin was completely converted into the cleaved form, but no high molecular mass furin could be detected because it had not reached the compartment where sialic acid is attached to the N-linked glycans. Endo H treatment (data not shown) proved that this cleaved form has reached the medial Golgi cisternae because no shift was detected. Although under these conditions the endogenous furin of CV-1 cells is present in the TGN, HA could not be activated during single expression because it had not reached this compartment and no or inadequate amounts of endogenous furin had been synthesized during the monensin incubation period. On the contrary, newly synthesized bovine furin had accumulated in the early and medial-Golgi region and cleaved the coexpressed HA into HA1 and HA2. Whereas HA1 produced in the absence of monensin migrated as a diffuse band (Fig. 6 B, bottom, third and forth lane), HA1 produced in the presence of the ionophore ran as a sharp band, proving that cleavage occurred before the compartment in which sialic acid is attached was reached. These results show that furin can process substrates in the early Golgi region even before it arrives in the TGN where it is temporarily concentrated.

**COOH-terminal Truncation of Furin Depends on Low pH and Ca2+ Ions**

Pulse-chase labeling (Fig. 2 C) strongly hinted at an intra-
with a furin-specific antiserum and the other one with an antiserum raised against FPV. After immunoprecipitation proteins were analysed by SDS-PAGE on 10% gels and fluorography. (B) 4.5 h after infection with recombinant vaccinia viruses encoding bovine furin (VV:bfur) or FPV hemagglutinin (VVHAvw) or coinfection with both of these viruses, CV-1 cells were labeled with 35S-methionine for 15 min and chased for 60 min with cold methionine. During starvation, labeling and chase period cells were either incubated in the presence or absence of brefeldin A (2 μg/ml). Cell lysates of the single infections were incubated with the respective antisera whereas lysates of coinfected cells were divided up into two halves one of which was incubated with the furin-specific and the other one with an antiserum raised against FPV. For further treatment of samples and for legends see A. Arrows on the left indicate positions of molecular mass standards. Arrows on the right mark the positions of precursor (HA) and cleavage products (HA1 and HA2) of the FPV hemagglutinin; ○, pro-furin; △, autocatalytically cleaved furin; ▽, autocatalytically cleaved furin with sialylated carbohydrate side chains.

cellular site of COOH-terminal truncation of furin, because cell lysates contained the smaller, 80–90-kD mature protein. Cleavage of the luminal domain could also be shown to be a late event taking place after acidic residues have been attached. Taken into consideration that furin predominantly localizes to the TGN, where a low pH is generated by proton pumps, the question arose whether acidotropic reagents, like ammonium chloride or chloroquine, might influence this proteolytic cleavage step. Furin labeled under normal conditions could only be precipitated from the conditioned medium (Fig. 7 A, lane S) after a 4-h chase period (Fig. 7 A, first and second lane), whereas in the presence of 50 mM ammonium chloride (Fig. 7 A, lanes labeled NH4Cl) most of the labeled furin was immunoprecipitated from the cell lysates (lane C) in the mature 104-kD form as could be confirmed by its resistance to endoglycosidase H (data not shown). Only small amounts of the truncated form were found in the supernatants (lane S). Almost complete inhibition of truncation was observed in the presence of 80 μM chloroquine (Fig. 7 A). Chloroquine not only raises the pH in the acidic compartments but has also an inhibiting effect on endogenous furin (Oda et al., 1986; Nagahama et al., 1991; Ohuchi et al., 1994). These data indicate that truncation of furin takes place in a compartment with an acidic milieu.

To further study if truncation depends on Ca2+ ions we tested the effect of the ionophore A23187 (Fig. 7, lanes A23187). Only high molecular mass furin was precipitated from cell lysates after the 4-h chase period, proving that Ca2+ ions are indispensable for the COOH-terminal truncation. In this case the cells were depleted of Ca2+ ions only after the 45-min labeling period because autocatalytic processing which is also Ca2+-dependent should occur during the labeling period. Otherwise the labeled, uncleaved pro-furin could not leave the ER (See Fig. 4 B). To prove that Ca2+ removal had not inhibited transport of the autocatalytically cleaved furin and that it had reached the trans-Golgi region, endo H digestions were performed (Fig. 7 B). Cleaved and transported furin represented most of the labeled furin because it was endo H resistant and shifted to 92 kD after deglycosylation with N-glycanase F. In summary, we conclude that truncation of furin is caused by a proteolytic reaction which takes place at low pH inside the cells. The Ca2+ dependence of this process might indicate that it is mediated by an autocatalytic activity, but it cannot be ruled out that other proteases are responsible for this reaction.

**Discussion**

A general feature of subtilisin and subtilisin-like proteases of eukaryotes is their synthesis as prepro-enzymes which are proteolytically processed by an intramolecular cleavage event removing a propeptide from the catalytic domain (Power et al., 1986; Brenner and Fuller, 1991; Leduc et al., 1992; Creemers et al., 1993). Whereas autoprocessing of subtilisin leads to secretion of the active protease, the eukaryotic enzymes kexin and furin remain temporarily mem-
COOH-terminal truncation of furin. Fluorograms of furin immunoprecipitated from cell lysates (C) and supernatants (S) of cells in which acidic compartments had been neutralized or whose intracellular Ca²⁺ concentrations had been reduced, 4.5 h after infection with recombinant vaccinia viruses encoding bovine furin (Vv-furin) cells were labeled for 45 min with [³⁵S]methionine and chased for 4 h with cold methionine. Supernatants and cell lysates were either incubated with NH₄Cl (50 mM) or chloroquine (80 μM) during starvation, labeling and chase period. To reduce intracellular Ca²⁺ concentrations cells were washed extensively with Ca²⁺ free medium after the labeling period and during chase they were incubated with Ca²⁺ ionophore A23187 carbohydrates were removed. In this respect compartmentalization of pro-furin conversion resembles maturation of the yeast protease kexin (Wilcox and Fuller, 1991) very closely and therefore this conservation of the intracellular location of propeptide removal seems to be typical for maturation of these subtilisin-like proteases in the secretory pathway of eukaryotic cells.

Ca²⁺ ions have been shown to be indispensable for substrate cleaving activity (Klenk et al., 1984; Bresnahan et al., 1990; Hatsuazawa et al., 1992; Molloy et al., 1992; Steineke-Grober et al., 1992). Our experiments employing the ionophore A23187 reveal that autocatalytic cleavage is strictly dependent on Ca²⁺ ions, too.

Figure 7. Acidotropic reagents or Ca²⁺ removal interfere with COOH-terminal truncation of furin. Fluorograms of furin immunoprecipitated from cell lysates (C) and supernatants (S) of cells in which acidic compartments had been neutralized or whose intracellular Ca²⁺ concentrations had been reduced, 4.5 h after infection with recombinant vaccinia viruses encoding bovine furin (Vv-furin) cells were labeled for 45 min with [³⁵S]methionine and chased for 4 h with cold methionine. Supernatants and cell lysates were either incubated with NH₄Cl (50 mM) or chloroquine (80 μM) during starvation, labeling and chase period. To reduce intracellular Ca²⁺ concentrations cells were washed extensively with Ca²⁺-free medium after the labeling period and during chase they were incubated with Ca²⁺ ionophore A23187 in medium lacking Ca²⁺ ions. (B) After immunoprecipitation of furin from cells treated with the Ca²⁺ ionophore A23187 carbohydrates were digested with endoglycosidase H or N-glycanase F to check if autocatalytic processing and transport of furin had occurred. The arrow on the left of fluorograms marks the position of the molecular mass standards; □/□, autocatalytically cleaved furin with sialylated carbohydrate side chains; ○, autocatalytically cleaved furin after truncation of the COOH-terminal membrane spanning domain; Δ, deglycosylated form of the protein.

Soon after translation of pro-furin, autoprocessing occurs with a half time of less than 15 min (Fig. 2 A). During the whole process, no conversion of the high mannose type N-glycans of pro-furin and cleaved furin occurs, indicating that the ER is the primary site of autoprocessing. Even after this process, the N-glycans of cleaved furin remain endo H sensitive for more than 30 min. The experiments employing brefeldin A underlined the important role the ER plays during these early events. The observation that in the presence of BFA, which prevents exit of proteins from the ER (for review see Klausner et al., 1992), profurin cleavage occurs (Fig. 6 A) indicates that no post ER compartment is involved in autocaltalysis. These results are in contrast to a previous report postulating that autoprocessing might be a late event. This was inferred from studies on a soluble furin mutant with an attached KDEL sequence that mediated ER retention (Rehemtulla et al., 1992). However, studies of Molloy et al. (1994) who used soluble, epitope tagged KDEL- and membrane-bound, epitope-tagged furin mutants and our data on unmutated bovine furin independently show that furin does not have to enter a post ER compartment for pro-furin propeptide removal. In this respect compartmentalization of pro-furin conversion resembles maturation of the yeast protease kexin (Wilcox and Fuller, 1991) very closely and therefore this conservation of the intracellular location of propeptide removal seems to be typical for maturation of these subtilisin-like proteases in the secretory pathway of eukaryotic cells.

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Mutants whose autocatalytic cleavage is severely impaired show that inhibition of autoprocessing leads to accumulation of pro-furin in the ER (Molloy et al., 1994; Creemers, J. W. M., M. Vey, W. Schäfer, T. A. Y. Ayoubi, A. J. M. Roebroek, H.-D. Klenk, W. Garten, and W. J. M. Van de Ven, manuscript submitted for publication). This observation suggests that propeptide removal might be a prerequisite for exit out of the ER. Our studies on wild-type furin reveal that carbohydrates of wild-type pro-furin remain endo H sensitive when autocaltalysis is impaired, for example in the absence of Ca²⁺ ions or in the presence of DTT (Fig. 5), hinting at an arrest of pro-furin in an early compartment. As a consequence, only correctly cleaved furin can leave the ER and enter the Golgi region. This means that furin has to pass a quality control. A similar regulatory function of the ER has been described for the influenza virus hemagglutinin. During folding, hemagglutinin adopts intermediate conformations which interact with the chaperone BiP (Gething et al., 1986). BiP binding is Ca²⁺ dependent and, in the absence of Ca²⁺ ions, formerly retained molecules enter the Golgi apparatus (Suzuki et al., 1991). Pro-furin does not seem to interact with BiP, because in the absence of Ca²⁺ ions it does not bear complex N-glycans, indicating that it is still in the ER (Fig. 4 B). The recently discovered chaperone calnexin has been shown to bind to glycoproteins like the heavy chain of MHC class I molecules (Galvin et al., 1992) and the complement factor C3 (Ou et al., 1993). Thus, calnexin, which is ubiquitously and abundantly expressed in the ER, might be the candidate chaperone mediating pro-furin retention.

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We further investigated the compartment of furin substrate cleavage. Previous studies showed that furin is concentrated in the Golgi compartment (Bresnahan et al., 1990; Misumi et al., 1991). Furthermore, it has been known for a long time that substrates are cleaved late after their synthesis (Klenk et al., 1974), and in recent subcellular fractionation experiments cleavage products of substrates produced by coexpressed furin could only be precipitated from Golgi fractions (Wasley et al., 1993), hinting at a late compartment of furin action. Furin mutants captured in the ER via a KDEL sequence were not able to process coexpressed substrates in this early compartment (Rehemtulla et al., 1992; Molloy et al., 1994). We coexpressed FPV hemagglutinin with wild-type furin in the presence of BFA and found that substrate cleavage did not occur (Fig. 6 A), proving that the ER and the intermediate salvage compartment do not allow substrate cleavage, whereas autoprocessing is not inhibited (Fig. 6 A). Thus, we conclude that furin is either blocked, maybe by binding to some kind of chaperone, or has to be further modified in order to gain substrate cleaving activity. In this respect furin resembles the kexin protease of *S. cerevisiae* because the yeast protease is not able to cleave substrates if it is captured in the ER (Julius et al., 1984b). Thus, furin and kexin seem to be not only related by a conserved structure of their catalytic domains but also their maturation in the secretory pathway seems to underly similarly conserved control mechanisms.

The next compartments furin enters are the cis- and medial-Golgi cisternae, and we asked whether furin can activate substrates there. Coexpression in the presence of monensin clearly showed that furin can cleave HA in the early Golgi region where no endogenous furin activity could be detected (Fig. 6 B). This finding demonstrates that furin gains substrate cleaving activity when it enters the Golgi region, even before it reaches the trans-Golgi cisternae and TGN where it is concentrated (Molloy et al., 1994; Schäfer et al., submitted). Therefore, the specific conditions in the TGN, including low pH and sugar modifications like sialylation, are not crucial for proteolytic activity. They may be of importance for another step in the intracellular lifecycle of furin (see Fig. 7 and Discussion below).

During transport to and concentration of furin in the TGN, the N-linked carbohydrate side chains of furin are processed and we have monitored these modifications by endoglycosidase digestions (Fig. 2). An earlier report on the sugar modifications of rat furin suggested that pro-furin and autocatalytically cleaved furin gain complex N-glycans (Misumi et al., 1991) very late after synthesis, which is not consistent with our results. In our studies, carbohydrates of pro-furin never showed endo H resistance. Even cleaved furin could hardly be detected in a 98-kD endo H-resistant form under normal conditions. Since rat furin was labeled late after onset of furin expression, whereas in our system furin was labeled early after onset of furin expression, differences in the assay systems may account for the different results. The endo H-resistant, cleaved furin that can be detected in our studies is sialylated and this modification leads to a significant shift in electrophoretic mobility (Figs. 2 and 3). Sialylation of furin proves that the protease has reached the trans-Golgi cisternae and the TGN, where sialyltransferases are located. Recently, evidence was provided that furin is not strictly retained in the TGN but it cycles between the plasma membrane and the TGN which implicates that furin appears in the endosomal compartment (Molloy et al., 1994) where it might activate exogenous substrates like the protective antigen of *B. anthracis* or exotoxin A of *Pseudomonas*. On the other hand, acidic compartments seem to be necessary for the final step of the intracellular lifecycle of furin which is discussed below. A schematic drawing (Fig. 8) illustrates the complex maturation pathway of furin.

Figure 8. Schematic drawing of the maturation pathway of furin. Compartmentalization of furin maturation. Prepro-furin is synthesized into the ER lumen, in which signal and propeptide are removed. After propeptide cleavage furin enters the Golgi region and gains substrate cleaving activity. In the TGN N-glycans acquire sialic acid and the protease is concentrated there. Furin then proceeds to the plasma membrane but is retrieved back to the TGN probably via the endocytic pathway (Molloy et al., 1994). Finally, a truncation process in an acidic compartment releases the enzyme from the membrane and furin is secreted from cells.

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Intracellular furin levels seem to be strictly controlled, because endogenous furin cannot be detected by immunocytochemical methods, and only catalytic amounts are present in the TGN, although furin mRNA seems to be continuously present in all cells tested so far. Other transcriptional or posttranscriptional regulatory mechanisms might be involved, but our experiments strongly hint at an important posttranslational control by a second proteolytic cleavage step that leads to secretion of an anchorless but active form. Truncation of furin, although less efficient than observed here, has been reported before (Wise et al., 1990; Rehemtulla et al., 1992; Vidricaire et al., 1993). Molloy et al. (1994) have recently reported efficient shedding of truncated furin from cells expressing wild type or a mutant that lacked the cytoplasmic tail. Our results with endogenous furin of MDBK cells now strongly suggest that truncation is a process by which endogenous furin is modified also under physiological conditions, because an 85-kD furin is the only furin-immunoreactive protein that can be detected in the enriched Golgi fractions. Earlier studies on this endogenous furin have shown that the electrophoretic mobilities of furin after elution from a molecular sieve column increased in subsequent fractions (Stieneke-Gröber et al., 1992) hinting at a similar modification of terminal sugars by sialylation which can be seen for mature, membrane bound and secreted bovine furin causing a heterogeneously migrating band in the SDS-PAGE (Figs. 2 C and 3). The studies on the overexpressed wild-type bovine furin further show that secretion of truncated furin is indeed a very efficient process by which cells lose this membrane-bound enzyme. After 4 h of chase (Fig. 2 B), the labeled protein completely disappeared from cells, and most of it was found in the media in a truncated and heavily sialylated form. In addition, the studies strongly hint at an intracellular site of cleavage because the COOH-terminally truncated furin could also be precipitated from cell lysates (Fig. 2). Furthermore, when we raised the pH of acidic compartments by addition of weak bases like chloroquine or ammonium chloride, truncation was inhibited. This observation suggests that concentration of furin in an acidic compartment like the TGN, on the one hand, might provide permanently low levels of active, intracellular furin by truncation of the membrane bound form. On the other hand, accumulation in the TGN does not interfere with cleavage activity because it occurs between pH 6 and 8 (Molloy et al., 1992; Stieneke-Gröber et al., 1992). With regard to substrate specificity low intracellular furin levels also make sense, because high levels of furin cause unspecific cleavage (Walker et al., 1993). So far no direct information on the enzyme(s) catalyzing this truncation reaction exist, but an earlier report (Rehemtulla et al., 1992) proposed an autocatalytic cleavage similar to propeptide processing. The restricted substrate specificities of propeptide processing (Leduc et al., 1992; Creemers et al., submitted) and substrate cleavage (Molloy et al., 1992; Nakayama et al., 1993; Watanabe et al., 1993) argue against an autocatalytic process, because no typical motif like R-X-K/R-R, R-X-X-X-X-K/R-R or R-X-X-K/R-R is present in the region where furin can be clipped off the membrane. Ca2+ dependence and inhibition by chloroquine, which both have been observed with substrate cleavage as well as truncation, do not provide direct evidence for an autocatalytic reaction, either. Identification of the exact peptide bond that is hydrolyzed and studied employing protease inhibitors might give some further information on the nature of the enzyme(s) involved.

In conclusion, our results and the recent studies on furin demonstrate that maturation of this TGN protease is a precisely tuned process, consisting of individual, successive steps which are performed in different compartments and depend on Ca2+ ions, oxidizing conditions and pH among other so far unidentified mechanisms.

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