Intracellular itinerary of internalised β-secretase, BACE1, and its potential impact on β-amyloid peptide biogenesis

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Abbreviations: Aβ, β-amyloid peptide; BACE1, beta site APP-cleaving enzyme; APP, amyloid precursor protein; TGN, trans-Golgi network

Synopsis

BACE1 cleavage of the amyloid precursor protein (APP) is the initial step in the formation of the amyloidogenic β-amyloid peptide. This paper reports that cell surface BACE1 is internalised by the AP2/clathrin dependent pathway and traffics to early endosomes then recycling endosomes. In contrast, internalised wild-type APP traffics to late endosomes/lysosomes. A BACE1/TGN38 chimera that recycles via the TGN is more efficient in β-amyloid production than wild-type BACE1 indicating that the recycling itinerary of BACE1 influences β-amyloid biogenesis.

Abstract:

β-secretase (BACE1) cleavage of the amyloid precursor protein (APP) represents the initial step in the formation of the Alzheimer’s disease associated amyloidogenic β-amyloid peptide. Substantive evidence indicates that APP processing by BACE1 is dependent on intracellular sorting of this enzyme. Nonetheless, knowledge of the intracellular trafficking pathway of internalised BACE1 remains in doubt. Here we show that cell surface BACE1 is rapidly internalised by the AP2/clathrin dependent pathway in transfected cells and traffics to early endosomes and Rab11-positive, juxtanuclear recycling endosomes, with very little transported to the TGN as has been previously suggested. Moreover, BACE1 is predominantly localised to the early and recycling endosome compartments in different cell types, including neuronal cells. In contrast, the majority of internalised wild-type APP traffics to late endosomes/lysosomes. To explore the relevance of the itinerary of BACE1 on APP processing, we generated a BACE1 chimera containing the cytoplasmic tail of TGN38 (BACE1/TGN38), which cycles between the cell surface and TGN in an AP2-dependent manner. Wild-type BACE1 is less efficient in β-amyloid production than the BACE1/TGN38 chimera, highlighting the relevance of the itinerary of BACE1 on APP processing. Overall the data suggests that internalised BACE1 and APP diverge at early endosomes and that β-amyloid biogenesis is regulated in part by the recycling itinerary of BACE1.

Key words: endosomal sorting, β-secretase, BACE1, recycling endosomes, trans-Golgi network, amyloid precursor protein, amyloid β peptide.
Introduction

Senile plaques, consisting primarily of deposits of β-amyloid peptide (Aβ), are a pathological hallmark of Alzheimer’s disease (AD) (1). Aβ is derived from sequential proteolysis of the amyloid precursor protein (APP) by β- and γ-secretases (2). β-cleavage of APP is first carried out by the membrane-anchored aspartyl protease BACE1 (3-7) to generate a membrane-bound fragment (C99) that is subsequently cleaved by γ-secretase to yield Aβ. An alternative, non-amyloidogenic pathway involves the cleavage of APP by α-secretase resulting in the release of the peptide p3 after γ-cleavage, a pathway that precludes Aβ production (8).

The relative abundance of proteolytically-processed APP products, including Aβ, is likely to reflect the intracellular localisation of APP and the different membrane secretases. The level of APP processing by BACE1 is dependent on the trafficking pathways of both BACE1 and APP, and also competition by α-secretase, which cleaves APP to promote the protective or the non-amyloidogenic pathway. As α-secretase is localised predominantly at the cell surface, it is likely that the majority of APP processing at the cell surface is along the non-amyloidogenic pathway (9, 10), whereas intracellular APP processing would predominantly involve the amyloidogenic pathway. As a member of the aspartyl protease family, BACE1 has an acidic pH optimum for activity and a number of intracellular location(s) for APP processing by BACE1 have been reported (11-13); the prevailing view is that Aβ is generated mainly in the endosomes and TGN (14-18). As γ-secretase is present in multiple intracellular compartments (13), the processing of APP by BACE1 is the critical step in the biogenesis of Aβ. The potential relevance of endosomal sorting in neurological disease is highlighted by recent findings suggesting that defects in membrane trafficking events of the early endosomes, as a consequence of depletion of retromer components or the sortilin-related receptor SorLA for example, may be an underlying cause promoting enhanced APP processing and Aβ biogenesis in AD (19, 20, 21).

Defining the intracellular trafficking pathways of BACE1 and APP is clearly important to understand the cell biology of Aβ production and to maximise opportunities for targeted inhibitors to BACE1. BACE1 is synthesised as a pro-protein in the endoplasmic reticulum (ER) where it is post-translationally modified by the addition of N-glycans and the subsequent removal of the prodomain as it traverses the Golgi and is transported to the cell surface (22-24). BACE1 then recycles between the cell surface and endosomes (11, 13). However, the precise trafficking pathways of BACE1 and APP are not well defined. Recent studies have reported that BACE1 is internalised by either an AP2-mediated clathrin-dependent pathway (17) or alternatively a clathrin-independent Arf6-dependent pathway (25). Moreover, some studies have indicated that internalised BACE1 may recycle via the early endosomes and TGN (26, 27), which represents a defined endosomal retrograde transport pathway (28). Other studies have suggested that BACE1 is internalised and recycled from endosomes to the plasma membrane (PM) (24) or transported from early to late endosomes/lysosomes for degradation (29). Likewise, newly synthesised APP is delivered to the cell surface and then rapidly internalised. Many (30-32), but not all (33), studies suggest that the internalised APP is either degraded in lysosomes and/or recycled back to the cell surface, most likely from early or late endosomes, whereas other studies have suggested that APP is recycled to the TGN (18). Hence there is considerable uncertainty about the trafficking pathways of both BACE1 and APP and very few studies have analysed both membrane proteins within the same system.

The trafficking itineraries of transmembrane proteins are dependent on sorting signals in their cytoplasmic tail. Sorting motifs in the cytoplasmic tail of BACE1 have been identified (reviewed in (34)), however, the identification of internalisation motifs does not provide predictions on the precise intracellular itinerary of cargo. More detailed knowledge of the BACE1 trafficking pathway is required, information which may provide potential avenues for sequestering BACE1 from APP and reducing the yield of the amyloidogenic Aβ peptide. Here we demonstrate that BACE1 recycles via the recycling endosomes, and not the TGN, in a variety of mammalian cells, including neurons. On the other hand internalised APP takes a different route and the majority is transported down the lysosomal pathway. Furthermore, we demonstrate that the itinerary of BACE1 is relevant to the regulation of Aβ production, as a BACE1 chimera that cycles via the TGN generates higher levels of Aβ levels compared with wild-type BACE1. Thus, we propose that the normal recycling of BACE1 through the recycling endosomes segregates BACE1 from APP and may be a factor that regulates the level of secreted Aβ in normal cells.

Results

Steady state distribution of BACE1 in cultured cells

There remains considerable uncertainty about the precise intracellular trafficking routes of BACE1 (34). Previous studies have indicated that BACE1 is internalised into early endosomes and recycles to the plasma membrane (PM) via early/late endosomes or the TGN (17, 26, 27, 29). To accurately define the intracellular location and trafficking itinerary of BACE1 we have used a set of well-defined organelle-specific markers for the early and late endosomes, recycling endosomes and Golgi/TGN. Initially we examined the steady state distribution of BACE1 in HeLa cells transfected with a BACE1 construct. Full-length BACE1 was associated with punctate structures scattered throughout the cytoplasm and concentrated in the perinuclear region of the cell (Fig. 1A). Substantial overlap of BACE1 was detected with the early endosome marker, EEA1, whereas in contrast there was very little overlap with the late endosome/lysosomal marker, CD63 (Fig. S1). Dual staining with the Golgi stack marker, GM130, (Fig. 1A) or with the TGN marker, golgin-97 (not shown) showed some but not extensive overlap with BACE1 suggesting...
that the majority of the juxtanuclear localised BACE1 may be located in a compartment distinct from the Golgi. Notably, the recycling endosome is also located in the juxtanuclear region; however, the possibility that BACE1 could be localised to the recycling endosome has not been carefully investigated. Indeed, by staining for endogenous Rab11 as a marker for the recycling endosome, BACE1 co-localised extensively with the recycling endosome in transfected HeLa cells (Fig. 1B). The use of GFP-Rab11 as a marker for recycling endosomes also demonstrated considerable overlap with BACE1 (not shown), confirming the specificity of the endogenous Rab11 staining. Quantification of the staining patterns revealed that, at steady state, BACE1 showed a 31.8% overlap with the early endosome marker, EEA1, 31.9% overlap with Rab11 and 4.4% overlap with the TGN marker, golgin-97, in HeLa cells (Fig. 1C).

To determine the relevance of these findings in HeLa cells, the steady-state distribution of BACE1 was then analysed in other cell lines including CHO cells, as this line allows excellent discrimination between recycling endosomes and other compartments and have been widely used to study APP processing events (35, 36), and SK-N-SH cells as this line represents a well-defined neuronal cell line relevant to AD (37). Significantly, very little overlap of BACE1 with the Golgi marker, GM130, was detected in either CHO or SK-N-SH cells (Fig. 1A) whereas extensive co-localisation of BACE1 and endogenous Rab11 was apparent in both CHO and SK-N-SH cells (Fig. 1B). These data clearly demonstrate that a substantial proportion of BACE1 is located at the juxtanuclear region, however, the possibility that BACE1 could localise to the recycling endosome has not been carefully investigated. Indeed, by staining for endogenous Rab11 as a marker for the recycling endosome, BACE1 co-localised extensively with the recycling endosome in transfected HeLa cells (Fig. 1B). The use of GFP-Rab11 as a marker for recycling endosomes also demonstrated considerable overlap with BACE1 (not shown), confirming the specificity of the endogenous Rab11 staining. Quantification of the staining patterns revealed that, at steady state, BACE1 showed a 31.8% overlap with the early endosome marker, EEA1, 31.9% overlap with Rab11 and 4.4% overlap with the TGN marker, golgin-97, in HeLa cells (Fig. 1C).

Intracellular itinerary of BACE1
To further define the intracellular itinerary of full-length BACE1, we tracked the transport of BACE1 from the PM to the Golgi apparatus using an antibody internalisation assay. HeLa cells were transfected with the wild-type BACE1 construct and incubated with anti-BACE1 antibodies on ice. As expected, the antibody-BACE1 complexes were restricted to the cell surface at 4°C (Fig. 2A, 0 min). Surface bound antibody-BACE1 complexes were then internalised at 37°C over a 60 min period. Antibody-BACE1 complexes were efficiently internalised; by 10-15 min the surface BACE1 had been internalised and was located in endosomal structures distributed in the cell periphery and throughout the cytoplasm (Fig. 2A, 15 min). There was considerable overlap of BACE1 with the early endosome marker EE1A at this early time point (Fig. 2B) as well as with endogenous Rab11 (Fig. 2E). After 60 min of internalisation, the majority of the internalised BACE1 remained located in punctate structures throughout the cytoplasm with a substantive co-localisation of internalised BACE1 with Rab11, particularly in the perinuclear region of the cell (Fig. 2C, E). Quantitative analysis showed an overlap of BACE1 with Rab11 of ~30% at either 15 or 60 min of 37°C chase. In contrast, there was very little co-localisation detected with the TGN marker, golgin-97 (Fig. 2A, 0 min), or the Golgi stack marker (GM130) (not shown), over the 60 min period (Fig. 2F).

The transferrin receptor (TfR) is known to recycle between the cell surface and the recycling endosomes (38). Given the intracellular transport of cell surface BACE1 to a Rab11-positive compartment, BACE1 may follow a similar intracellular trafficking route as that used by TfR. Therefore, we tracked BACE1 together with transferrin (Tf). Substantial co-localisation of Tf and BACE1 was observed at 15 min and 60 min after internalisation. At 60 min internalisation a significant level of staining was located in the perinuclear region of the cell (Fig. 2D), consistent with the location of recycling endosomes. Quantitative analyses showed that the overlap between the internalised Tf and BACE1 was 45.0±1.9% at 15 min and 42.1±1.15% at 60 min internalisation (Fig. 2G). Hence, BACE1 and TfR appear to have a similar intracellular trafficking route, further confirming that BACE1 localises to the recycling endosomes.

To determine whether Rab11 is required for the intracellular transport of BACE1 we examined the effect of the dominant-negative mutant Rab11S25N on the endosomal trafficking of BACE1. Rab11S25N has previously been shown to affect the transport of cargo sorted from the early/recycling endosomal compartments (39). HeLa cells were transfected with either GFP-Rab11wt or GFP-Rab11S25N together with BACE1 and an internalisation assay for BACE1 performed. We compared the transport of BACE1 through the EEA1-positive early endosome compartment in Rab11wt and mutant Rab11S25N over-expressing cells. After 15 min internalisation, approximately 35% of BACE1 co-localised with EEA1 in both Rab11wt and Rab11S25N expressing cells (Fig. S2). Whereas the level of BACE1 in the EEA1-positive compartment then decreased in Rab11wt expressing cells over a subsequent period of 45-75 min, no decrease was observed in the mutant GFP-Rab11S25N over-expressing cells (Fig. S2), indicating a block in transport from early endosomes and suggesting that a functional Rab11 is required for BACE1 intracellular trafficking.

Our previous studies have demonstrated that the membrane protein, TGN38, is transported directly from the early endosomes to the TGN (40). To determine the point of divergence of BACE1 and TGN38 along the endocytic pathway we investigated the trafficking of BACE1 and TGN38 simultaneously. Dual antibody uptake assays showed extensive co-localisation of BACE1 and TGN38 in punctate structures after 15 min internalisation (Fig. S3), which are likely to represent early endosomes based on the above time course study of BACE1. Internalisation for 45 min or 90 min resulted in segregation of BACE1 and TGN38, with internalised TGN38 localised predominantly with a TGN marker and BACE1 localised to independent peripheral punctate structures (Fig S3). These data suggest that the divergence of BACE1 and TGN38 most likely occurs from the early endosomes. Collectively these data indicate that BACE1 in internalised and is transported from early endosomes to recycling endosomes, similar to the trafficking of the TfR.

BACE1 internalisation is mediated by AP2 and clathrin
There have been contradictory reports regarding the internalisation pathway of BACE1 from the PM to early endosomes. The di-leucine motif in the cytoplasmic tail of BACE1 has been suggested to facilitate clathrin-mediated endocytosis (reviewed in (34)); recent reports have concluded that BACE1 is endocytosed predominantly either by an AP2-dependent (17) or an AP2-independent, Arf6-dependent (25) pathway. To assess the relevance of AP2 and clathrin in the endocytosis of BACE1, we have directly determined the machinery requirements for the internalisation of BACE1 from the cell surface. A characteristic of the clathrin-mediated pathway is very rapid internalisation of cargo. To assess the impact of clathrin depletion on BACE1 distribution, the ubiquitous heavy chain isofrom of clathrin (CHC17) was silenced in HeLa cells with a previously defined human CHC17 siRNA target sequence (41) followed by transfection with wild-type BACE1. Immunoblotting showed that the level of clathrin heavy chain was significantly reduced (~80%) in CHC17 siRNA transfected cells (Fig. 3A). As expected TIR accumulated on the surface of CHC17-depleted cells (Fig. 3B). The depletion of clathrin heavy chain also resulted in a dramatic reduction in the internalisation of BACE1 from the cell surface over a 15 min period (Fig. 3C). We also verified using the FACS-based internalisation assay above that BACE1 endocytosis was dependent on the di-leucine motif in the cytoplasmic tail. As expected mutation of the di-leucine motif in the cytoplasmic tail of BACE1 (BACE1LL/AA) resulted in an enhanced level of BACE1 at the cell surface of the transfected cell population (not shown). Cell surface BACE1LL/AA was not readily internalised as demonstrated by no observed change in the percentage of cell surface BACE1-positive cells after 10 min internalisation at 37°C (Fig. 3C).

Given that the RNAi experiments did not completely block the internalisation of BACE1, which could be due to incomplete silencing, we also inhibited clathrin-mediated endocytosis with the selective clathrin inhibitor, pitstop 2 (42) and assessed the internalisation of BACE1 (Fig. 4). To quantitate internalisation, cell surface proteins of intact cells were biotinylated with Sulfo-NHS-SS-biotin and the levels of biotinylated BACE1 assessed by immunoblotting of biotinylated complexes pulled out on NeutrAvidin beads. The selective release of the cell surface biotinylated molecules with the cell-impermeable reducing agent glutathione (GSH) allows the discrimination of cell surface BACE1 from internalised BACE1 (Fig. 4A). After 30 min or 60 min internalisation at 37°C, a substantial proportion of biotinylated BACE1 was protected from glutathione release, indicating internalisation (Fig. 4B). At 30 min >50% of biotinylated BACE1 was protected from the cell impermeable glutathione, whereas at 60 min at 37°C, following glutathione treatment at 30 min and 60 min, ~40% of biotinylated BACE1 was protected; the reduction between 30 and 60 min of biotinylated BACE1 suggests recycling of internalised BACE1 back to the cell surface. Incubation with pitstop 2 reduced the level of protected BACE1 and the majority of the biotinylated BACE1 (~80%) remained susceptible to cell surface release by glutathione (Fig 4C, D). On the other hand incubation with pitstop 2 followed by a 1 h washout on ice, and followed by an incubation at 37°C for 30 min, resulted in ~50% of biotinylated BACE1 being protected from glutathione reduction (Fig. 4C, D). Using antibodies to label cell surface BACE1, confocal microscopy confirmed that BACE1 was retained on the cell surface in pitstop 2-treated cells, whereas cell surface antibody-BACE1 complexes were internalised into EEA1-positive structures following washout of the inhibitor (Fig. 4E magnified image). These results show that the clathrin inhibitor, pitstop 2, blocks internalisation of the majority of surface BACE1 and that the effects of pitstop are reversible.

Therefore, collectively, these data confirm the requirement of the LL motif for internalisation of BACE1, and demonstrate that BACE1 is internalised predominantly via the AP2/clathrin-mediated pathway.

**Alteration of the intracellular distribution of BACE1**

The above data indicates that cell surface BACE1 is endocytosed by an AP2/clathrin dependent pathway and subsequently recycles via the recycling endosomes and not the TGN. To further discriminate between the recycling endosomes and the TGN, and to ascertain if there was functional consequence of re-directing BACE1
trafficking to the TGN, we constructed a BACE1 mutant which is predicted to cycle via the TGN, by swapping the cytoplasmic tail of BACE1 with that from a membrane protein which undergoes retrograde transport from endosomes to the TGN. Specifically, we replaced the 23-residue cytoplasmic tail of BACE1 with the 34-residue cytoplasmic tail of the membrane protein TGN38 (BACE/TGN38, Fig. 5A). TGN38 is known to recycle between the cell surface and the TGN via early endosomes, a trafficking route dependent on the cytoplasmic tail (40, 43). Both BACE1 and TGN38 are type I transmembrane proteins with C-terminal cytoplasmic tails. Firstly, we assessed the internalisation pathway of the BACE/TGN38 chimera. BACE/TGN38 was internalised from the cell surface by the AP2/clathrin pathway of the BACE/TGN38 chimera. BACE/TGN38 cytoplasmic tails. Firstly, we assessed the internalisation pathway of the BACE/TGN38 chimera. BACE/TGN38 was internalised from the cell surface by the AP2/clathrin pathway as expected; silencing CHC17 or AP2 μ2 resulted in the inhibition of internalisation of cell surface BACE/TGN38 (not shown). Therefore the initial internalisation pathway of wild-type BACE1 and the BACE/TGN38 chimera are similar.

In contrast to wild-type BACE1, at steady state a considerable proportion of BACE/TGN38 was located at the Golgi of transiently transfected HeLa cells (Fig. 5B). Furthermore, antibody uptake assays showed that BACE/TGN38 was efficiently transported from the cell surface via endosomes to the Golgi over a 60 min period (Fig. 5C, D); notably at the 60 min time-point there was substantial (~50%) overlap with the Golgi marker, GM130 (Fig. 5C, D) indicating efficient retrograde transport of BACE/TGN38 to the Golgi.

These data demonstrate that the itinerary of the BACE/TGN38 chimera differs from wild-type BACE1, and that, in contrast to wt BACE1, the chimera is transported from endosomes to the TGN. Collectively, the data also provides further discrimination in the trafficking of cargo to either the recycling endosomes or the Golgi.

**Internalised APP predominantly traffics along the lysosomal pathway**

A number of studies have shown that APP is internalised from the PM to early endosomes and to the lysosomes for degradation (30-32). However, other studies suggest that APP can be recycled to the TGN (18) and moreover it is also possible that the transmembrane domain may influence endosomal sorting events and trafficking. To ascertain whether APP has sorting signals in the cytoplasmic tail to direct this membrane cargo to the lysosomes, we have generated a cleavage-resistant APP chimeric protein containing the transmembrane domain and cytoplasmic tail of APP with the luminal reporter domain, CD8, a widely used reporter (45). Cell surface CD8/APP was labeled with anti-CD8 antibodies and complexes were internalised at 37°C; after 30-60 min, internalisation antibody-CD8(APP complexes showed co-localisation with both GFP-Rab7(Q67L) and the endogenous late endosome/lysosome marker LAMP1 marker (Fig. S4A, B). Quantitation showed there was a progressive increase in the co-localisation with GFP-Rab7(Q67L) and LAMP1 over a 60 min period (Fig. S4D, E), with only low levels of co-localisation with the TGN marker, p230 (Fig. S4F). Taken together these results confirm that the majority APP is internalised to late endosomes/lysosomes, suggesting that the intracellular transport pathways of BACE1 and APP differ.

**Effect of altered trafficking of BACE1 on APP processing**

To determine if there was a functional consequence of redirecting BACE1 trafficking from the recycling endosomes to the TGN, we compared the efficiency of Aβ biogenesis by wild-type BACE1 and the BACE/TGN38 chimera. For these experiments we employed a CHO cell line stably expressing APP695wt (35) as we have established that CHO-APP cell lines provide a sensitive system for the detection of Aβ. CHO-APP stable clones expressing the BACE constructs were generated. Clones were selected which expressed very similar levels of BACE1 and BACE/TGN38; an immunoblot showed that BACE was present in the BACE1 and BACE/TGN38 CHO clones in a ratio of 1:1.25 respectively, based on normalisation with α-tubulin (Fig. 8A). An immunoblot of a dilution series of the lysates showed a consistent ratio of BACE at various dilutions (not shown). The BACE/TGN38 chimera co-localised extensively with the Golgi marker, GM130, under steady-state conditions, whereas very little wild-type BACE1 co-localised with GM130 in the stable cell internalisation (Fig. 6A) or for up to 60 min (Fig. 6F). Likewise minimal co-localisation was detected between internalised APP and the TGN marker, p230/golgin245 (Fig. 6G); the latter was chosen as the TGN marker due to compatibility of the different antibodies used in the dual staining. Conversely, antibody-APP complexes co-localised extensively with Rab7-positive structures for extensive periods (Fig. 6C) with ~30% of APP co-localising with GFP-Rab7(Q67L) after 60 min internalisation (Fig. 6E). The constitutive active from of Rab7 was used to demark the late endosomes as GFP-wt Rab7 shows only a low level of membrane staining with the majority of the fusion protein located in the cytosol.

One complication with the use of wild-type APP is that the tracking antibody recognizes the luminal domain of APP and therefore does not recognize the cleaved, membrane-associated, forms of APP. The intracellular sorting signals of APP are located in the cytoplasmic tail, and based on our recent work on endosomal sorting (44) it is also possible that the transmembrane domain may influence endosomal sorting events and trafficking. To ascertain whether APP has sorting signals in the cytoplasmic tail to direct this membrane cargo to the lysosomes, we have generated a cleavage-resistant APP chimeric protein containing the transmembrane domain and cytoplasmic tail of APP with the luminal reporter domain, CD8, a widely used reporter (45). Cell surface CD8/APP was labeled with anti-CD8 antibodies and complexes were internalised at 37°C; after 30-60 min, internalisation antibody-CD8(APP complexes showed co-localisation with both GFP-Rab7(Q67L) and the endogenous late endosome/lysosome marker LAMP1 marker (Fig. S4A, B). Quantitation showed there was a progressive increase in the co-localisation with GFP-Rab7(Q67L) and LAMP1 over a 60 min period (Fig. S4D, E), with only low levels of co-localisation with the TGN marker, p230 (Fig. S4F). Taken together these results confirm that the majority APP is internalised to late endosomes/lysosomes, suggesting that the intracellular transport pathways of BACE1 and APP differ.
lines (Fig. 7A). The availability of Golgi markers for CHO cells is more restricted than HeLa cells, however, the extensive co-localisation of BACE/TGN38 with the Golgi stack marker GM130 clearly demonstrates Golgi localisation. Internalisation experiments were also performed to verify that the trafficking of BACE1 in these stable CHO-APP clones was similar to that of transiently transfected cells. As only low levels of surface expression were present in these stable clones, we incubated the CHO clones with anti-BACE1 antibody at 37°C to allow a continuous uptake of BACE1 from the cell surface. After 2 h internalisation, there were substantial levels of antibody-BACE/TGN38 co-localised with GM130, whereas very little internalised antibody-wild-type BACE1 was detected in the Golgi region (Fig. 7B). Quantitation revealed a 2.3-fold higher level of internalised BACE/TGN38 at the Golgi compared with wild-type BACE1 (Fig. 7C). Therefore, the stable CHO-APP clones expressing the BACE constructs behaved in a similar fashion to the transiently transfected cells.

To determine the levels of Aβ production in the BACE1 stable CHO-APP lines, we analysed the secreted products for the presence of Aβ. The level of the secreted product of α-secretase (sAPPα) was also determined as a measure of overall secretion. Whereas similar levels of sAPPα was detected from BACE1 and BACE/TGN38 CHO-APP cells, there was a 3-fold higher level of Aβ in the supernatants from the BACE/TGN38 CHO-APP cells compared with wt BACE1 CHO-APP cells (Fig. 8). The levels of Aβ in these experiments were normalised for the level of BACE1 protein in each clone (Fig. 8A). A 3-fold difference in Aβ production between the CHO clones was very reproducible and observed in four independent experiments (average of four experiments was 2.8-fold) with two independent clones of the constructs (Fig. 8C). Taken together, these data demonstrate that the precise intracellular trafficking pathways of BACE1 influence the yield of Aβ generated and moreover show that recycling of BACE1 through the recycling endosomes reduces the yield of Aβ compared with recycling via the TGN.

Discussion

The aggregation of Aβ to form toxic oligomers and amyloid plaques is considered to be central to the pathogenesis of Alzheimer’s disease, and the cleavage of APP by BACE1 is the initial step in the generation of Aβ peptide (46). Substantive evidence indicates that the processing of APP by BACE1 is regulated by the intracellular sorting and trafficking of the enzyme, hence an understanding of membrane transport of BACE1 is of critical importance for the development of therapeutics to treat AD. BACE1 recycles between the plasma membrane and a perinuclear compartment, previously considered to be the TGN (26, 27). The identity of this perinuclear compartment is highly relevant to the appreciation of potential major sites of APP processing. Our study has revealed that BACE1 is internalised by AP2/clathrin to early endosomes and then traffics from the early endosomes to the juxtanuclear recycling endosome (in both non-neuronal and neuronal cell lines), and not to the TGN/Golgi as previously thought. In contrast APP is internalised and transported predominantly along the endosomal/lysosomal pathway. Our work has also revealed that the trafficking route of BACE1 is pertinent to APP processing and Aβ generation. A BACE/TGN38 chimera, which is efficiently transported to the TGN by virtue of a defined TGN-retrieval motif in the TGN38 cytoplasmic tail (47), generates Aβ levels considerably higher than wild-type BACE1. Given our findings, and the comparative itineraries of APP and BACE1 in transfected cells, we propose that internalised BACE1 and APP diverge at the early endosomes and that the transport of BACE1 to the recycling endosomes may be an important factor in regulating Aβ production.

The internalisation of BACE1 from the cell surface has been attributed to a di-leucine motif (17, 24-26). The di-leucine motif in the cytoplasmic tail of BACE1 is part of a more complex motif known as the acidic cluster-di-leucine motif (DXXLL), composed of DISLL residues, (24, 26). Mutation of the leucines to alanines was previously shown to decrease internalisation of BACE1 (24, 48), thus implicating the DISLL sequence as an internalisation motif. Here we have confirmed the importance of the LL residues on internalisation of BACE1 and demonstrated that rapid internalisation is mediated by AP2 and clathrin endocytosis as silencing clathrin heavy chain, AP2 μ2 or AP2 α-adapin resulted in the inhibition of internalisation of BACE1. The effect on BACE1 internalisation by inhibition of clathrin-mediated endocytosis with the small molecule inhibitor, pitstop 2, is also consistent with this conclusion, although the specificity of pitstop 2 for clathrin-mediated endocytosis has very recently been questioned (49). While this manuscript was in preparation another report was published which also demonstrated that endocytosis of BACE1 is mediated by AP2 and furthermore showed that the DDTEGR sequence fits the consensus AP2 binding motif (17). Structural studies have recently revealed an interaction between the acidic cluster di-leucine motifs with the hydrophobic pockets on the σ2 subunit of AP2 (50), providing a molecular basis for clathrin-mediated endocytosis of BACE1. Another recent report had suggested that BACE1 endocytosis was independent of clathrin and AP2 but rather dependent on Arf6 (25). We were unable to demonstrate that Arf6 silencing had any effect on the rapid internalisation of BACE1 but did result in a modest increase in the level of BACE1 accumulating in early endosomes (unpublished data), findings consistent with the known role of Arf6 in intracellular recycling (51). However, our findings do not exclude a role for Arf6 in the slow internalisation of BACE1 by non-clathrin pathways.

BACE1 has a long half-life and recycles between endosomes and the PM multiple times during its lifespan (24). The perinuclear recycling endosome, also known as the endocytic recycling compartment (ERC), was distinguished from other endosomes and the Golgi by the specific marker Rab11 and by the localisation of internalised TIR (52), and the exclusion of EEA1 and Golgi markers. Moreover, a role for Rab11 and recycling endosomes in BACE1 trafficking is supported by the finding that Rab11(S25N) perturbs intracellular transport of BACE1. Our findings suggest that BACE1 has a similar intracellular trafficking pathway as TIR to the recycling endosomes. The perinuclear recycling
endosome is often centrally located, juxtaposed alongside the microtubule organising centre, and in close proximity to the Golgi (53, 54). The perinuclear recycling endosome is characterised by the presence of Rab11 and has a tubular morphology from which transport carriers destined for the cell surface are derived (reviewed in (38, 55)). The observation that BACE1 is internalised to early endosomes and then to the Rab11-positive compartment suggests that a substantial level of internalised BACE1 takes the slow recycling route via the recycling endosomes back to the PM, a suggestion compatible with a role of Arf6 in recycling. A ‘fast recycling route’ from the early endosomes back to the PM has also been documented for TR (55), and it is possible that some BACE1 may also use the fast recycling route.

Our study used transfected cells to examine the intracellular distribution of BACE1 at steady state conditions and the intracellular itinerary after internalisation from the cell surface. It is possible that the level of exogenous BACE1 may influence its’ intracellular trafficking itinerary. However, we believe this is unlikely for the following reasons: Firstly, BACE1 was not retained in the ER indicating newly synthesized BACE1 was folded efficiently and was transported out of the ER. Secondly, the majority of BACE1 in transfected cells was located in early endosomes and recycling endosomes and very little BACE1 was detected in late endosomes (CD63 marker) at steady state in transfected cells. Moreover cell surface BACE1 was efficiently transported to the recycling endosomes, a pathway considered to be signal dependent (56). If over expression was influencing trafficking then one would expect inefficient sorting in the early endosomes resulting in default delivery to the late endosome and lysosome. The observation that there is efficient sorting and transport of BACE1 to the recycling endosome in transiently transfected HeLa cells and stable CHO cells argues that the expression levels have not saturated the machinery for sorting. Other studies have also used exogenous expression of BACE1 to monitor internalisation as surface levels of endogenous BACE1 are very difficult to detect (17, 27, 57).

In contrast to BACE1, APP is internalised from the PM to early endosomes and is transported to lysosomes for degradation (30-32). Our studies here have confirmed these findings and, furthermore have demonstrated that APP is not transported to the recycling endosomes. Use of a reporter construct CD8/APC allowed the intracellular itinerary of the cytoplasmic tail of APP to be dissected without the complication of intracellular processing events for full-length APP. CD8/APC was internalised also to the late endosomes/lysosomes, confirming that the trafficking signals of APP direct the bulk of this membrane protein to lysosomes, at least in transfected HeLa cells.

A large proportion of APP is found at early endosomes at steady state and indeed the bulk of Aβ production is thought to occur within this compartment (34), consistent with the preference of the aspartyl protease activity of BACE1 for an acidic pH. Evidence that the early endosome is a major site for Aβ biogenesis is supported by recent findings that the endosomal coat complex, retromer, is involved in the trafficking of BACE1. Significantly, RNAi silencing of one of the retromer components, Vps35, results in elevated levels of Aβ in cultured cells (19, 21). Retromer defects have also been detected in the brains of AD cases, notably the downregulation of Vps35 (19). In addition, analysis of brains of mice expressing reduced levels of retromer (VPS26+/−) has revealed elevated levels of Aβ and associated hippocampal dysfunction (20). Recent studies have indicated that retromer may regulate multiple transport steps from the early endosome, rather than solely facilitating retrograde transport. For example, retromer has been shown to be required for recycling of the β2-adrenergic receptor directly back to the cell surface from early endosomes (58), and for sorting and transport from the early endosome to the recycling endosomes in addition to the well established role in endosome-to-TGN transport (40, 59); hence the potential importance of retromer on BACE1 trafficking.

Interestingly, SorLA, a receptor of the Vps10-domain family, binds APP and is itself sorted in a retromer-dependent manner (60-62), and moreover perturbations in SorLA levels manifests changes in trafficking of APP and Aβ production. Sortilin, another member of the Vps10 family, also influences the endosomal trafficking of APP (63). These findings highlight the potential relevance of endosomal sorting and cargo receptors in BACE1 and APP trafficking and in neurodegenerative disease.

We assessed the impact of redirecting BACE1 via the TGN rather than the recycling endosomes. One advantage of directly manipulating the trafficking of BACE1, compared with the silencing of endosomal machinery components, is that the normal trafficking of APP remains unaffected. Redirecting BACE1 via the TGN rather than recycling endosomes resulted in a ~3-fold increase in Aβ production. Previous studies have shown that Aβ production can occur in the secretory pathway (64, 65), and the enhanced production of Aβ by BACE/TGN38 is likely to result from the TGN as a potential site for APP cleavage, in addition to early endosomes. The TGN has an acidic pH which is favourable for BACE1 activity, and recycling of BACE/TGN38 via the TGN will expose newly synthesised APP to high levels of BACE1 activity. Hence the enhanced production of Aβ may result from BACE1 processing of newly synthesised APP arriving in the TGN. We did not detect any difference in the ratio of Aβ1-40/42 products between wild-type BACE1 and BACE/TGN38 (unpublished data). In addition to Asp1 β-site cleavage to generate Aβ1-40/42, BACE1 can also cleave at Glu11 (β'-site cleavage) which results in truncated Aβ11-40/42 products. Previous studies have indicated that there may be enhanced APP β-site cleavage by BACE1 in the TGN compared with other locations (66), although the physiological relevance of these truncated Aβ peptides remains unclear. Further work is clearly required to define the pathway for Aβ production by the BACE/TGN38 chimera. Nonetheless, our finding of a 3-fold increase in Aβ1-40/42 production by BACE/TGN38 construct compared with wild-type BACE1 is likely to represent a physiologically significant difference in Aβ production. Downregulation of the retromer component Vps26 results in 2-3 fold elevated levels of Aβ and associated hippocampal dysfunction (67), illustrating the significance of a 3-fold difference with regards to in vivo cytotoxicity.
In summary, the individual trafficking itineraries of APP and BACE1 are critical for the processing of APP along the amyloidogenic pathway resulting in Aβ generation. We have demonstrated that cell surface APP and BACE1 have distinct trafficking itineraries. Importantly, we have demonstrated that BACE1 is transported predominantly to the recycling endosome and not the TGN as previously proposed, whereas APP is transported to late endosomes/lysosomes. We propose that internalised BACE1 and APP are segregated at the early endosomes by independent sorting events, and that BACE1 is predominantly transported to the recycling endosomes for transport back to the PM (Fig. 9). Through the segregation of BACE1 at the early endosome for recycling back to the cell surface via the recycling endosomes, internalised BACE1 is prevented from reaching the TGN. Thus the normal trafficking of BACE1 may be protective against excessive Aβ production. These findings have important implications for the development of therapeutics, and could provide a novel approach to inhibit Aβ production by interfering with BACE1 recycling.

Materials and Methods

Plasmids, antibodies and reagents

pCDNA4/TO BACE1 was a gift from Michael Cater, Mental Health Research Institute, Victoria, Australia. GFP-Rab11wt, GFP-Rab11(S25N) and GFP-Rab7(67L) are N-terminus fusion constructs with GFP, as described (68). TGN38-CFP encodes a C-terminal fusion protein with the fluorescent CFP (69). mCherry-FYVE\_EAA\(_1\) was generated by subcloning the region encoding the FYVE domain of EEA1 from GFP-FYVE\_EAA\(_1\) (70) into the BGII and EcoRI sites of the mCherry-N1 vector (gift from Dr Roger Tsien UCSD, CA, USA). Mouse monoclonal anti-β amyloid precursor protein was from Invitrogen (Camarillo, CA, USA). The full length APP cDNA in the pIRESpuro2 expression vector (Clontech Laboratories Inc., Mountain View, CA, USA) has been described (71). Mouse antibodies to clathrin heavy chain (clone 23), Rab11 (clone 47), EEA1 (clone 14), GM130 (clone 35) and TGN38 (clone 2/TGN38) were purchased from BD Bioscience (NSW, Australia). Mouse monoclonal antibodies to human CD63 (clone mx-49.129.5) was from Santa Cruz. Mouse monoclonal antibodies to golgin-97 (A-21270) were purchased from Life Technologies (NY, USA). Mouse monoclonal antibodies specific for Aβ5-8 (W02) have been described (72). Mouse monoclonal antibodies to AP2 α-adapin (clone 100/2) and Arf6 (clone 2A-1) were from Sigma Aldrich (USA) and Santa Cruz Biotechnology, Inc (CA, USA) respectively. Mouse monoclonal antibodies to human CD9a (clone PA-T8) were from eBioscience (CA, USA). Affinity purified rabbit polyclonal anti-BACE1 (EE-17) was purchased from Sigma Aldrich (USA). Rabbit polyclonal antibodies to the μ2 subunit of AP2 and LAMP1 were purchased from Epitomics Inc. (CA, USA) and Abcam (Cambridge, UK) respectively. Human autoantibodies to p230 have been described previously (73).

Conjugates used for immunofluorescence were goat anti-rabbit IgG-Alexa Fluor 568, goat anti-rabbit IgG Alexa Fluor 488, goat anti-mouse IgG-Alexa Fluor 488, goat anti-human IgG Alexa Fluor 647 and goat anti-human IgG Alexa Fluor 568 from Molecular Probes (Invitrogen, Carlsbad, California, USA). Horseradish peroxidase-conjugated sheep anti-rabbit IgG and anti-mouse IgG were from DAKO Corporation (Carpinteria, California, USA). Can Get Signal Immunoreaction Enhancer Solution was purchased from Toyobo Life Science Department (Osaka, Japan).

Generation of BACE/TGN38 and CD8/APP chimeric constructs

A unique AgeI site was introduced at the junction of the transmembrane and cytoplasmic domains in the nucleotide sequence of wild-type BACE1 using the site-directed mutagenesis using the primers 5' CACCTGTCCTATGGTGTGACGGG TGCGCTCGAGTCTAGAGGCCCC 3' and 5' GGCCTCTCTGACCTGAGCGGACC GGTACACACATGGAAGCAAGTGAGTCGATGCT 3' to produce pCDNA4/TO BACE1 AgeI. The cytoplasmic tail sequence of rat TGN38 was amplified with flanking AgeI restriction sites by PCR using the primers 5' GGGGGACCGGGTCACAACAA ACGCAAA 3’ and 5’ CCCCCACCGTTCACATTTGAACTTGAAGTTCGATGCT 3’. The amplified product was purified and digested with AgeI before ligation into AgeI-digested pCDNA4/TO BACE1 AgeI. The pCDNA4/TO BACE1/LL/AA mutant was obtained using the primers 5' TTTCGTCATGACATCTCCGCTGTAATTGAGATGAC TCGAGTCTGAG 3' and 5' AGACTCCGAGTCATCATTGACGCGAGATGTC ATCAGCAAA 3' by site-directed mutagenesis, using pCDNA4/TO BACE1 as a template.

To generate the CD8-APP construct, the transmembrane and cytoplasmic domains of APP was amplified with flanking 5’HindIII and 3’ Xba sites by PCR using the primers CCAAGCTTGGTGCAATCATTGGACTCAT3' and TCTAGAGGGGTCACAACA ACGCAAA 3’. The CD8 lacuneral domain was isolated from the CD8- M6PR vector construct (74) by PCR and including flanking 5’XhoI and 3’ HindIII sites by PCR using the primers 5'TGCTCTAGACTAGTTCTGCATCTGCTCAA3'. Materials and Methods

Cells and transient transfections

HeLa cells and human neuronal SK-N-SH cells were maintained as semi-confluent monolayers in DMEM supplemented with 10% (v/v) fetal calf serum (FCS), 2 mM L-glutamine, 100 units/μl penicillin and 0.1% (w/v) streptomycin (C-DMEM). Chinese Hamster Ovary (CHO) cells were grown in RPMI media supplemented with 10% (v/v) fetal calf serum (FCS), 2 mM L-glutamine, 100 units/μl penicillin and 0.1% (w/v) streptomycin (C-RPMI). Cells were maintained in a humidified 10% CO₂ atmosphere at 37°C. For transient transfections, HeLa cells were seeded as monolayers and transfected using Fugene 6 (Roche Diagnostic, Basel, Switzerland) and for CHO and SK-N-SH cells, cells were transfected with Lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) and Opti-MEM according to manufacturer’s instructions. Transfections were carried out in C-DMEM or C-RPMI at 37°C, 10% CO₂ for 24 h.
RNA interference
The human CHC17(1) specific siRNA duplex is as described in (41). The human μ2(1), α-adaptin(1) and CHC17(2) specific siRNA duplexes are described by (75), siRNA duplexes for Arf6(1) (76), Arf6(2) (77), μ2(2) (78) and α-adaptin(2) (17) are as described. All above described siRNA target sequences were purchased from Sigma Proligos (Lismore, Australia).

Generation of stable cell lines
A CHO cell line stably expressing APP695WT (CHO-APP) (35) was maintained in C-RPMI with 7.5 μg/ml of puromycin (Sigma Aldrich). BACE1-CHO-APP and BACE/TGN38-CHO-APP cell lines were generated by transfecting the CHO-APP cell line with either wild-type BACE1 or BACE/TGN38 constructs respectively, using Lipofectamine 2000 (Invitrogen, Carlsbad, California, USA). Stably expressing cells were selected and maintained in C-RPMI medium with 7.5 μg/ml puromycin and 50 μg/ml zeomycin (Sigma Aldrich). Monoclonal cell lines were generated by limiting dilution.

Indirect immunofluorescence
Cells on coverslips were fixed with 4% paraformaldehyde for 15 min, followed by quenching in 50 mM NH₄Cl/PBS for 10 min. Cells were permeabilised in 0.1% Triton X-100/PBS for 4 min and blocked in 5% FCS/PBS for 20 min to reduce non-specific binding. Monolayers were incubated with primary and secondary conjugates as described (79) and confocal microscopy performed using a Leica TCS SP2 imaging system. For endogenous Rab11 staining, cells were fixed with 10% trichloroacetic acid in PBS for 15 min, followed by quenching with 30 mM glycine/PBS. Cells were then permeabilised and with 0.1% Triton X-100/PBS and blocked with 5% FCS/PBS as described above. Monolayers were incubated with mouse anti-Rab11 antibodies diluted in Can Get Signal Solution A (Toyobo, Japan), followed by incubation in secondary conjugates diluted in blocking solution. Images were collected independently for multi-colour labelling.

Internalisation assay
Cells were transfected 24 hr prior to the internalisation assay. Rabbit polyclonal BACE1 antibodies (2 μg/ml), APP antibodies (10 μg/ml) or CD8 antibodies (2.5 μg/ml) were incubated with cells on ice for 30 min, cells washed, and the BACE1 or APP antibody-bound complexes normalised to the levels of cell associated tubulin. BACE1 or antibody-BACE/TGN38 complexes was determined by taking the sum of overlapping pixels between BACE1 and respective markers, divided by the total number of BACE1 pixels within each cell. Data obtained from fluorescence intensity of Golgi-localised antibody-BACE1 or antibody-BACE/TGN38 complexes was expressed as the mean, +/- SEM and analysed by an unpaired, two-tailed, student t-test. A p<0.05 (*) was considered significant, p<0.01(**) highly significant and p<0.001 (***)) very highly significant. The absence of a p value indicates the differences were not significant.

Immunoblotting
For immunoblotting of cell lysates, cell extracts were dissolved in reducing sample buffer and samples resolved on a 4-12% NuPAGE gradient gel and immunoblotting was performed as described (74). Blots were imaged using the Gel Pro Analyzer program (MediaCybernetics, Bethesda, USA) and levels of protein normalised to the levels of cell associated tubulin.

Cell surface biotinylation
HeLa cells were transfected with BACE1 for 24 hours and cell surface biotinylation was performed as described (17). Briefly, cells were incubated in 1 mM Sulfo-NHS-SS-Biotin (Pierce, Rockford, IL, USA) in PBS for 1 h on ice. Unreacted ester was quenched in cold 50 mM Tris-HCl for 10 min. Cells were then lysed in cell lysis buffer (50mM Tris-HCl, pH 7.5, 150 mM NaCl, 2 mM EDTA, 1% (v/v) Triton X-100, containing protease inhibitors (Roche Applied Science, Indianapolis, IN, USA). For internalisations, cells were surface biotinylated as described, and then incubated at 37°C for 30 min or 60 min. Cells were then treated with cleavage buffer (50 mM glutathione (Sigma Aldrich), 90 mM NaCl, 1.25 mM CaCl₂, 1.25 mM MgSO₄, 0.2% BSA pH 8.6) for 1 h on ice, followed by treatment with 50 mM iodoacetamide for 30 min to quench remaining glutathione. Cells were lysed in cell lysis buffer in the presence of 2.5 mM iodoacetamide. For pitstop treatment, surface biotinylated cells were treated with 30 μM pitstop 2 (Abcam, Cambridge, UK) in PBS for 15 min on ice prior to incubation at 37°C for 30 min, with or without a 1 h washout with cold D-MEM, followed by glutathione cleavage. Lysates were extracted with cell lysis buffer. All lysates were incubated with NeutrAvidin-agarose (Pierce) in a rotary shaker at 4°C for 2 hours. Beads were washed 3 times with cell lysis buffer and analysed by SDS-PAGE followed by immunoblotting with BACE1 antibody. Densitometry of bands was carried out using ImageJ.

FACS analysis of BACE1 internalisation
HeLa cells were transfected with BACE1 or BACE1LL/AA for 24 h and antibody-BACE1 uptake assay was carried out in suspension. For siRNA-treated cells, cells were first transfected with siRNA for 48 h followed by transfection with BACE1 for a further 24 h, prior to the BACE1 uptake assay. Monolayers were harvested by trypsin digest at 37°C for 3 min. Trypsin treatment had no effect on the levels of surface BACE1, as determined by comparison with EDTA harvested monolayers. After incubation at 37°C, cells were fixed with 4% PFA/PBS. Quenching and blocking was performed as described above, without permeabilisation. Cells were then stained with rabbit Alexa568-conjugated

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IgG and cells resuspended in FACS buffer (2 mM EDTA/PBS) for analysis on a Becton Dickinson LSRFortessa flow cytometer. Data was analysed with BD FACSDiva software (BD Biosciences).

**Analysis of secreted Aβ CHO cells**

Aβ blots of conditioned media from cell lines were carried out as described (35). Briefly, BACE APP CHO cells were seeded at 1.5 x 10^5 cells per well in a 12 well plate and grown to 90% confluency in C-RPMI supplemented with 50 μg/ml zeomycin and 7.5 μg/ml puromycin. The following day, selection media was removed and cells were washed twice with PBS. A 300 μl aliquot of C-RPMI without antibiotics was added to each well and after 16 h incubation media from each well was removed for Aβ analysis. Monolayers from corresponding wells were collected, cells lysed in reducing sample buffer and the samples analysis by immunoblotting to determine BACE expression levels. Conditioned media was centrifuged to pellet debris and 10 μl of conditioned media added to 10 μl of 2 x reducing sample buffer and 15 μl of the mixture was loaded per well of a 10% Tricine pre-cast gel for immunoblotting.

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**References**


Figure Legends

Figure 1. The steady state distribution of BACE1 in different cell types
HeLa, CHO and neuronal SK-N-SH cells were transfected with BACE1 for 24 h. Monolayers were fixed, permeabilised and stained with polyclonal BACE1 antibodies (red) and monoclonal antibodies to (A) GM130 (green) or (B) Rab11 (green). (C) HeLa cells, transfected with BACE1 as above were stained with mouse monoclonal antibodies to golgin-97 (TGN marker), Rab11 or EEA1 and the percentages of BACE1 pixels that overlapped with each marker were determined using the plugin OBCOL on the ImageJ program. Data is expressed as the mean +/- SEM (n=15). Stars indicate location of the nucleus. Bars represent 10 μm except for magnified images where bars represent 5 μm except for magnified images where bars represent 10 μm.
Figure 2. Internalised BACE1 transits the juxtanuclear recycling endosomes

(A) HeLa cells were transfected with BACE1 for 24 h and monolayers incubated with polyclonal BACE1 antibodies for 30 min on ice. Cells were washed in PBS and incubated in serum-free media for up to 60 min at 37°C, then fixed, permeabilised and stained for internalised antibody-BACE1 complexes with Alexa568-conjugated anti-rabbit IgG (red) and monoclonal golgin-97 antibodies (green). (B) HeLa cells were transfected with BACE1 for 24 h prior to the antibody internalisation assay, followed by staining with Alexa568-conjugated anti-rabbit IgG (red) and EEA1 antibodies (green). (C) HeLa cells were transfected with BACE1 for 24 h prior to the antibody internalisation assay, followed by staining with Alexa568-conjugated anti-rabbit IgG (red) and endogenous Rab11 (green). (D) The internalisation assay was carried out by incubating cells with both polyclonal BACE1 antibody and 568-conjugated transferrin (red), followed by staining with Alexa488-conjugated anti-rabbit IgG (green). The percentage of BACE1 at either (E) recycling endosomes or (F) the TGN was determined by calculating the percentage of total BACE1 pixels that overlapped with Rab11 or golgin-97 respectively using the plugin OBCOL on ImageJ program. (G) The percentage of total BACE1 pixels which overlapped with internalised 568-Tf. Data is expressed as the mean +/- SEM (n=20 for each time-point). Bars represent 10 μm, except for magnified images where bar represents 5 μm. In B the boxed region of the merge is magnified on the RHS.
Figure 3. Cell surface BACE1 is internalised via clathrin-mediated endocytosis
HeLa cells were transfected with either control siRNA, CHC17(1) siRNA, μ2(1) siRNA, AP2α aptin(1) siRNA or Arf6(1) siRNA for 48 h and transfected again with BACE1 for a further 24 h. (A) For immunoblotting, cells were lysed in SDS-PAGE reducing buffer and cell extracts were subjected to SDS-PAGE on 4-12% gradient polyacrylamide gel. Proteins were transferred to a PVDF membrane and probed with mouse anti-α tubulin antibody and mouse anti-CHC antibody, rabbit anti-μ2 antibody, mouse anti-α adaptin antibody or mouse anti-Arf6 antibody, using a chemiluminescence detection system. (B) For internalisation assays, cells were incubated with rabbit anti-BACE1 antibody and mouse anti-TfR antibody (OKT9) on ice for 30 min. Cells were then washed with cold PBS and incubated in serum-free media at 37°C for 15 min, and then fixed and permeabilised. Monolayers were then stained for antibody-bound BACE1 and antibody-bound TfR with Alexa568 conjugated anti-rabbit IgG (grey) and Alexa488 conjugated anti-mouse IgG respectively (grey). (C) HeLa cells were transfected with either BACE1 or BACE1LL/AA for 24 h. Monolayers were first lifted by trypsin digest at 37°C for 3 min. Cells were washed with PBS and incubated with rabbit anti-BACE1 antibody for 30 min on ice. Cells for the 0 min time-point were fixed immediately, whereas cells for internalisation were incubated in warm serum-free media for 10 min at 37°C, and then fixed. Cells were quenched in 50 mM NH4Cl and blocked in 5% FCS/PBS, without permeabilisation, and then resuspended in flow cytometry buffer and stained for cell surface BACE1 with Alexa568-conjugated IgG. 20,000 events were collected per sample and percentage of Alexa568-positive events indicated. Untransfected cells were incubated with primary and secondary antibodies to determine the gating, as indicated in the upper panel. Bar represents 10 μm.
A

B

C

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**Figure 4. Reversible inhibition of clathrin-mediated internalisation of BACE1**

(A) Cartoon depicting cell surface biotinylation of BACE1 and release of the biotin tag with GSH cleavage buffer. (B) HeLa cells were transfected with BACE1 for 24 h and cell surface biotinylation was carried out at 4°C. Biotinylated cells were either analysed directly (0 min –GSH), or subjected to glutathione (GSH) cleavage at 4°C (0 min +GSH); incubated in warm serum-free media at 37°C for 30 min, followed by cleavage (30 min +GSH); or incubated in serum-free media at 37°C for 60 min, with cleavage at both the 30 min and 60 min intervals (60 min +GSH). Cells were lysed and extracts were incubated with NeutrAvidin-agarose. Bound proteins were subjected to SDS-PAGE on a 10% polyacrylamide gel. Proteins were transferred to a PVDF membrane and probed with rabbit anti-BACE1 antibody using a chemiluminescence detection system. (C) Cell surface biotinylation and pitstop 2 treatment. Cell surface biotinylation was carried out on BACE1-transfected cells as described above. Cells were then incubated for 15 min with 30 μM pitstop 2 at 4°C and either incubated in serum-free media at 37°C for 30 min, followed by glutathione cleavage or subjected to a washout in cold C-DMEM on ice for 1 h, followed by internalisation at 37°C for 30 min and glutathione cleavage. Cells were lysed and processed as for B. (D) Bar graph of intensity of biotin bands after cell surface biotinylation and pitstop 2 treatment. Densitometric quantitation of bands was performed on 2 independent experiments using ImageJ. Values are represented as mean ± standard error. (E) HeLa cells were transfected with BACE1 for 24 h and monolayers were incubated with rabbit anti-BACE1 antibody for 30 min on ice. Cells were fixed or treated with 30 μM pitstop 2 for 15 min on ice. Pitstop 2-treated cells were subjected to internalisation at 37°C in serum-free media for 30 min, with or without a 1-hour washout in cold C-DMEM. Cells were fixed and permeabilised, followed by staining with Alexa568-conjugated anti-rabbit IgG. In the magnified image of the pitstop/washout sample, cells were stained with Alexa568-conjugated anti-rabbit IgG (red) and mouse anti-EEA1 antibodies (green). Data is from two independent experiments. Bars represents 10 μm.
A

4°C not cleaved

4°C cleaved

30min/37°C cleaved

B

Biotinylation pull down

Cell lysate

BACE1

C

Biotinylation pull down

Cell lysate

BACE1

D

Inhibition of BACE1 internalisation by pitstop 2

% of total biotinylated

BACE1

Time at 37°C (min) 0 0 30 30

GSH - + + +

Pitstop - - + +/-washout

E

0 min at 37°C

30 min at 37°C + pitstop

30 min at 37°C + pitstop/washout

BACE1

Magnified

BACE/EEA1

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Figure 5. The steady state distribution of BACE/TGN38 in HeLa cells
(A) Schematic of BACE1 and BACE/TGN38 constructs. (B) HeLa cells were transfected with BACE/TGN38 for 24 h and fixed, quenched and permeabilised, followed by staining with polyclonal BACE1 antibodies (red) and monoclonal GM130 antibodies (green). (C) HeLa cells were transfected with BACE/TGN38 for 24 h and monolayers were incubated with polyclonal BACE1 antibodies for 30 min on ice. Cells were washed in PBS and incubated in serum-free media for up to 60 min at 37°C, then fixed, permeabilised and stained for internalised antibody-BACE1 complexes with Alexa568-conjugated anti-rabbit IgG (red) and monoclonal GM130 antibodies (green) as the Golgi marker. (D) The proportion of BACE/TGN38 at the Golgi was determined by calculating the percentage of total BACE1 pixels that overlapped with GM130, using the plugin OBCOL on ImageJ. Data is expressed as the mean +/- SEM (n=20 for each time-point). Bars represent 10 μm and 5 μm (magnified images).
Figure 6. Internalised APP is transported to the late endosome and not the recycling endosomes

HeLa cells were transfected with APP and either (A) GFP-Rab11wt, (B) Cherry-FYVE or (C) GFP-Rab7(Q67L) for 24 h and monolayers incubated with monoclonal mouse APP antibodies for 30 min on ice. Cells were washed in PBS and incubated in serum-free media for up to 90 min at 37°C, then fixed, permeabilised and stained for internalised antibody-APP complexes with (A,C) Alexa568-conjugated anti-mouse IgG (red) or (B) Alexa488 conjugated anti-mouse IgG. The proportion of internalised APP at (D) early endosomes, (E) late endosomes, (F) recycling endosomes and (G) the TGN was determined by calculating the percentage of total internalised APP pixels that overlapped with EEA1, GFP-Rab7(Q67L), GFP-Rab11wt or p230 respectively, using the plugin OBCOL on ImageJ. Data is expressed as the mean +/- SEM (n=20 for each time-point). Bars represent 10 μm except for magnified images of B and C where bar represents 5 μm.
Figure 7. BACE1 internalisation in BACE1 and BACE/TGN38 CHO-APP stable cell lines

(A) Steady state distribution of BACE1 constructs in stable CHO APP cell lines. Stably-expressing CHO APP BACE1 or CHO APP BACE/TGN38 cells were fixed, quenched and permeabilised, followed by staining with BACE1 polyclonal antibodies (red), monoclonal GM130 antibodies (green) and DAPI (blue). (B) For BACE1 internalisation, CHO APP BACE1 or BACE/TGN38 cells were incubated with polyclonal BACE1 antibody in serum-free media at 37°C for 2 h. Monolayers were fixed, quenched and permeabilised, followed by staining with Alexa568-conjugated anti-rabbit IgG (red), monoclonal GM130 antibodies (green) and DAPI (blue). (C) The proportion of internalised antibody-BACE1 or antibody-BACE/TGN38 at the Golgi was determined by calculating the percentage of total BACE1 pixels that overlapped with GM130, using the plugin OBCOL on ImageJ program. Data is expressed as the mean +/- SEM (n=20 for each time-point). Bars represent 10 μm. *** indicates p<0.001.
A

BACE1  GM130  Merge
CHO BACE1

BACE/TGN38  GM130  Merge
CHO BACE/TGN38

B

Continuous 2h BACE1 Ab internalisation in stable CHO cell lines

BACE1  GM130  Merge
CHO BACE1

BACE/TGN38  GM130  Merge
CHO BACE/TGN38

C

% of BACE1 that overlaps with GM130

BACE1  BACE/TGN38

***

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Figure 8. Aβ levels from CHO APP BACE1 and BACE/TGN38 stable cell lines

(A) CHO-APP BACE1 or CHO-APP BACE/TGN38 cell lines were subjected to SDS-PAGE, transferred onto membrane and probed were probed with polyclonal BACE1 antibody to determine levels of BACE1 or BACE/TGN38. The membrane then stripped and reprobed with mouse anti-α-tubulin antibodies. 

(B) Conditioned media from stably-expressing CHO-APP BACE1 or CHO-APP BACE/TGN38 cell lines were subjected to SDS-PAGE, transferred onto membrane and probed with mouse W02 antibody to detect sAPPα and Aβ bands.

(C) Bar graph of the intensity of Aβ bands from four replicates. The levels of Aβ in these experiments were normalised for the level of BACE protein in each experiment. Densitometry of bands was carried out using ImageJ. * indicates p<0.05.
Figure 9. Model of trafficking itineraries of BACE1 and APP
(Left) Newly-synthesized BACE1 and APP are transported from the TGN to the PM. (Right) The pathways of internalised BACE1 and APP diverge at the early endosomes. Our data suggests that BACE1 is transported from the early endosome to the recycling endosomes, not the TGN, for return to the PM. On the other hand the bulk of APP is retained in the maturing early endosome for delivery to the late endosomes/lysosomes.
Table 1. Analysis of Internalisation of surface BACE1 by flow cytometry

HeLa cells were transfected with siRNAs for 48 h and transfected again with BACE1 for a further 24 h. Cell suspensions were incubated with rabbit anti-BACE1 antibody for 30 min on ice. Cells were fixed immediately or incubated in warm serum-free media for 10 min at 37°C. Cells were quenched and stained for cell surface BACE1 with Alexa568-conjugated IgG and analysed by flow cytometry. 20,000 events were collected per sample and levels of surface staining at 0 min and 10 min compared. Data presented for each treatment is the mean +/- SD from 3-4 experiments using two independent siRNA for each target.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>% Knockdown</th>
<th>% Reduction of BACE1-positive cells after 10 min internalisation</th>
</tr>
</thead>
<tbody>
<tr>
<td>control siRNA</td>
<td>NA</td>
<td>30.73 ± 5.94</td>
</tr>
<tr>
<td>CHC17 siRNA</td>
<td>80</td>
<td>11.58 ± 6.46</td>
</tr>
<tr>
<td>μ2 siRNA</td>
<td>75</td>
<td>10.58 ± 6.00</td>
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<tr>
<td>α-adaptin siRNA</td>
<td>85</td>
<td>12.15 ± 4.47</td>
</tr>
<tr>
<td>Arf6 siRNA</td>
<td>95</td>
<td>25.32 ± 6.37</td>
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</table>
Supplementary Figures

Figure S1  The steady state distribution of BACE1 in HeLa cells
HeLa cells were transfected with BACE1 for 24 h. Monolayers were fixed, permeabilised and stained with polyclonal BACE1 antibodies (red) and monoclonal antibodies to EEA1 (green) or CD63 (green), as indicated. Bar represents 10 µm.

Figure S2  Overexpression of Rab11S25 alters the transport of internalised BACE1
(A) HeLa cells were transfected with BACE1 together with either Rab11wt or Rab11S25N for 24 h and monolayers incubated with polyclonal BACE1 antibodies for 30 min on ice. Cells were washed in PBS and incubated in serum-free media for up to 90 min at 37°C, then fixed, permeabilised and stained for internalised antibody-BACE1 complexes with Alexa568-conjugated anti-rabbit IgG (red) and EEA1 antibodies (green). The percentage of BACE1 at the early endosome was determined by calculating the percentage of total BACE1 pixels that overlapped with EEA1 using the plugin OBCOL on ImageJ program. Data is expressed as the mean +/- SEM (n=12-15 for each time-point). ** indicates p<0.01.

Figure S3 Simultaneous trafficking of Internalised BACE1 and TGN38
(A) HeLa cells were co-transfected with BACE1 and TGN38-CFP for 24 h and transfected monolayers incubated with polyclonal rabbit anti-BACE1 antibodies and mouse anti-TGN38 antibodies for 30 min on ice. Cells were washed in PBS and incubated in serum-free media for up to 60 min at 37°C, then fixed, permeabilised and stained for internalised antibody-BACE1 complexes with Alexa568-conjugated anti-rabbit IgG (red) and for internalised antibody-TGN38-CFP complexes with Alexa488-conjugated anti-mouse IgG (green). The steady state localisation of TGN38-CFP was used as a marker of the TGN. In the top panel the boxed region of the merge is magnified on the RHS. Bars represent 10 µm.

Figure S4  Internalised CD8/APP is delivered to the late endosomes/lysosomes.
HeLa cells were transfected with (A) CD8/APP and GFP-Rab7(Q67L) or (B,C) CD8/APP alone for 24 h. Monolayers were incubated with mouse anti-CD8 antibodies on ice for 30 min, washed with cold PBS and incubated in serum-free media at 37°C for up to 180 min. Cells were stained with Alexa568-conjugated anti-mouse IgG and (B) rabbit anti-LAMP1 antibodies (green) and (C) human p230 antibodies (green). The percentage of CD8/APP pixels that overlapped with (D) GFP-Rab7, (E) LAMP1 or (F) p230 was determined using the plugin OBCOL on ImageJ. Data is expressed as the mean +/- SEM (n=15). Bars represent 10 µm and in magnified images, bars represent 5 µm.
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