RESEARCH COMMUNICATION

C8, a new member of the convertase family

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A novel subtilisin-like protein, PC8, was identified by PCR using degenerate primers to conserved amino acid residues in the catalytic region of members of the prohormone convertase family. PC8 was predicted to be 785 residues long and was structurally related to the mammalian convertases furin, PACE4, PC1 and PC2, sharing more than 50% amino acid identity over the catalytic region with these family members. PC8 possessed the catalytically important Asp, His, Asn and Ser amino acids, the homo B domain of this family of enzymes and a C-terminal hydrophobic sequence indicative of a transmembrane domain. Structurally, PC8 is more related to furin and PACE4 than to PC1 or PC2. Like furin and PACE4, PC8 mRNA was found to be widely expressed; this is in contrast with PC1 and PC2, which have a restricted distribution. Two transcripts, of 4.5 and 3.5 kb, were detected in both human cell lines and rat tissues. Unlike furin and PACE4, both of which map to chromosome 15, PC8 maps to chromosome 11q23–11q24, suggesting that this gene may have resulted from an ancient gene duplication event from either furin or PACE4, or conversely that these genes arose from PC8.

INTRODUCTION

Many biologically active proteins are synthesized as precursors which require specific endoproteolytic cleavage, commonly after dibasic residues, to generate their mature active forms. Over the last decade, a family of enzymes which is responsible for cleavage of pro-proteins to their biologically active forms has been identified [1–3]. This family of enzymes, known as ‘prohormone convertases’, are calcium-dependent serine proteinases exhibiting considerable sequence similarity to the bacterial subtilisins.

Kex2 was the first convertase to be described and was isolated and cloned from Saccharomyces cerevisiae [4,5]. Subsequently a further six members of this family, which exhibit highly conserved and similar catalytic domains, have been cloned from mammalian cells. These are furin [6,7], PC1 [8], also called ‘PC3’ [9], PACE4 [10,11], also identified as ‘PC7’ [12], PC2 [8,13], PC4 [14,15] and PC5 [16,17] also known as PC6 [18].

Furin, PACE4 and PC5 are expressed in a variety of tissues and cell lines. However, there are differences in their relative abundance [7,10–12,17–19]. Whereas PC1 and PC2 have more restricted distribution and are predominant in the anterior and intermediate pituitary respectively [8,20–22], PC4 is restricted to the testis [14,15].

The consensus motif for precursor cleavage by furin is marked by an R-X-K/R-R sequence. However, furin has also been shown to cleave at R-X-K-R and R-X-X-K/R-R sequences [1–3,19,23]. In contrast, PC1 and PC2 have been shown to direct precursor cleavage following dibasic acid amino pairs [20,24]; PC1 can also cleave at mono-arginyl sites with basic residues at the -4 or -6 position [25]. These sites are found in many pro-peptides and polypeptide precursors, and the mammalian pro-hormone convertases have been found to cleave a number of these, including pro-opiomelanocortin [24], proinsulin [26], pro-insulin receptor [27], prorelaxin [28], proalbumin [29], pro-(von Willebrand factor) [19] and pro-(β-nerve growth factor) [23].

In the course of studies aimed at understanding the post-translational processing of parathyroid hormone-related protein, which elicits multiple biological actions and contains a number of potential sites for convertase action, including two furin target sequences [30,31], we have identified a new member of the convertase family. Reported here is the identification, tissue distribution and chromosomal assignment of this new convertase which we have designated ‘PC8’.

MATERIALS AND METHODS

Cell culture

The human squamous cancer cell lines BEN, COLO 16 and the spontaneously immortalized human keratinocyte cell line HaCaT were cultured as previously described [31–33].

PCR oligonucleotides

oPC8 5′-GGGCCCCCA(G/A)CTNGC(G/A)CT(G/A)TA-3′, antisense strand oligonucleotide to amino acids 250–256 of human furin [7], oPC10 5′-TANCC(A/G)TC(A/G)CA(A/G)T(T/C/T)(A/G)CA-3′, antisense strand oligonucleotide to amino acids 303–308 of human furin [7], oPC12 5′-TGCTGCAT(A/G)TCNGCCA-3′, antisense strand oligonucleotide to amino acids 390–395 of human furin [7], oPC13

Abbreviations used: RT, reverse transcription (or transcribed); GAPDH, glyceraldehyde-3-phosphate dehydrogenase; UTR, untranslated region; PTHrP, parathyroid-hormone-related protein.

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Northern-blot analysis and reverse transcription PCR (RT-PCR)

Total RNA was extracted using guanidinium chloride as described [35] and polyadenylated RNA was recovered from total RNA using a QuickPrep Micro mRNA purification kit (Pharmacia Biotech Inc., Uppsala, Sweden). Northern analysis and hybridizations were performed as previously reported [33]. Specifically bound probe was quantified by phosphorImager analysis (Molecular Dynamics, Sunnyvale, CA, U.S.A.).

RT-PCR was performed on 5 µg total RNA isolated from cells or tissues as previously described [33,34]. PCR cycling conditions were: 94 °C for 30 s, 45–55 °C for 60 s (depending on the oligonucleotide combinations), 72 °C for 30 s to 2 min, for 40 cycles followed by 10 min elongation time at 72 °C. PCR products resolved on a 2 % (w/v)-agarose gel were subsequently extracted using Qiagen DNA purification kit (Qiagen Inc., Chatsworth, CA, U.S.A.) or Wizard PCR Preps (Promega Corp., Madison, WI, U.S.A.) or pGEMT (Pharmacia Biotech Inc.). DNA sequence analysis from both strands was performed using Sequenase sequencing (USB, Cleveland, OH, U.S.A.). Sequences were then compared with sequences in the GenBank™ databank.

Chromosomal localization

Metaphases were obtained from phytohaemagglutinin-stimulated lymphocytes from a healthy donor after thymidine synchronization and bromodeoxyuridine incorporation. The full-length cDNA HE6WCR40 was labelled with biotinyl-16-dUTP by random priming. Fluorescence in situ hybridization was performed as previously described [37].

RESULTS AND DISCUSSION

In order to identify convertase family members which may be involved in post-translational processing of parathyroid-hormone-related protein (PTHrP), we characterized PTHrP-producing cells for their convertase mRNA complement. Degenerate oligonucleotide primers (oPC2, oPC8, oPC10, oPC12 and oPC13) were designed to amino acid residues conserved between the catalytic regions of furin, PC1 and PC2. These primers, when used in RT-PCR, would allow the identification of known convertases and potentially new members of the convertase family. Total RNA from BEN cells was Reverse-transcribed with random hexamers, and 30 cycles of PCR undertaken with the primers oPC8 and oPC13. Resultant PCR fragments of approx. 190 bp were subcloned into pCRScript and sequenced. By this approach, 18 clones showing sequence similarity to furin, two to PACE4 and two to PC2, were recovered. In addition, another clone, BC20, of 187 bp was detected, and this was highly related to, but distinct from, the other convertase cDNA clones.

The cDNA sequence of BC20 was extended over the catalytic domain by RT-PCR with a BC20-specific sense-strand oligonucleotide, oPC33, and antisense-strand degenerate primers (oPC10 and oPC12) to conserved amino acids in the catalytic domain between furin, PC1 and PC2, which resulted in fragments 345 and 617 bp respectively; the sequences of these fragments have been previously reported [38,39]. It was clear that BC20 formed a new member of the mammalian prohormone convertase family and we refer to it now as ‘PC8’, adopting the nomenclature of Seidah et al. [8]. 5′ Sequences were amplified by anchored PCR from a human liver cDNA library in pcDNAIneo (Stratagene) using the PC8-specific antisense oligonucleotide (oPC34) and a sense-strand-specific primer to the T7 RNA polymerase promoter. Combined, these strategies yielded overlapping sequences from the 5′ untranslated regions (UTR) to the end of the catalytic domain of PC8 (Figure 1, nucleotides 1–1320). During database searches a cdNA clone, HE6WCR40 (GenBank™ accession no. H58761), representing an N-terminally truncated PC8 cDNA, was identified. This clone provided confirmation of the sequence obtained by RT-PCR and the complete 3′ coding region of PC8. Subsequently, the entire coding region of PC8 (GenBank™ accession no. U40623) was amplified from BEN cell RNA by RT-PCR and the sequence was identical with that determined for HE6WCR40. The cDNA clones predicted PC8 to be encoded by a mRNA species of 3.3 kb, composed of a 21 bp 5′ UTR, an open reading frame of 2355 bp coding for a protein of 785 amino acids and a 3′ UTR of 925 nucleotides. The 3′ UTR contained a polyadenylation signal, AAATAA, which was followed 12 nucleotides later by a poly(A) stretch (not shown). The primary protein sequence predicts a 42-residue signal peptide at the N-terminus, six potential N-linked glycosylation sites (residues 167, 175, 200, 241, 511 and 763) and a 22-amino-acid transmembrane region between residues 666 and 687 (Figure 1). The Met residue at position 36 is within a consensus Kozak sequence [40]. However, if this was the initiating AUG codon, the protein would lack a signal peptide. The N-terminus of mature PC8 is assumed to begin at Ser-142, following the sequence R-A-K-R, which fits the R-X-K/R-R consensus motif cleavage site for furin. While this manuscript was in preparation, a cDNA clone, HSU33849, identical with PC8 (except for a 131-nucleotide extension of the 5′ UTR and the last nucleotide of HSU33849 corresponding to nucleotide 2559 in Figure 1) and a partial cDNA clone, 45900, sharing structural similarity with PC8, were deposited in the GenBank™ database (accession nos. U33849 and H09374 respectively).

The chromosomal location of the PC8 gene was determined by in situ hybridization using the HE6WCR40 cDNA clone. A total of 30 metaphase preparations were analysed, and a specific labelling of bands 11q23–q24 was observed on four (seven cells), three (four cells), two (12 cells) and one (four cells) chromatid. No secondary peak was detected.

Comparison of PC8 with other mammalian convertases

An amino acid alignment of PC8 to the known human convertases furin, PACE4, PC1 and PC2 is given in Figure 3. Each possesses a signal peptide, an N-terminal domain, a catalytic domain, a homo B domain and a C-terminal domain. PC8 shares its highest identity with furin (54 %), PACE4 (53 %), PC1 (53 %) and PC2 (52 %) within the 289 amino acid catalytic domain. In addition, PC8 possesses the conserved amino acids Asp (197), His (228), Asn (329; Asp in PC2) and Ser (406), which are catalytically important. Outside the catalytic domain, the con-
Figure 1  Nucleotide and predicted amino acid sequences of PC8

The putative cleavage site to release the mature protein from the propeptide is indicated by the arrow ( ), and the putative transmembrane region is underlined. The subtilisin-like catalytic domain is boxed, and the catalytically important Asp, His, Asn, and Ser residues are indicated by . Consensus sites for Asn-linked glycosylation are marked by the open circles (D).

Figure 2  Idiogram of chromosome 11

Idiogram of the chromosome 11 homologue showing the unequivocal distribution of fluorescent spots on the 11q23–q24 bands for 15 metaphases.

Expression of PC8 mRNA

PC8 mRNA transcripts of 3.5 and 4.5 kb were found to be expressed in human cell lines (Figure 4A) and in all rat tissues examined (results not shown); these cell lines and tissues were also found to express mRNA for furin and PACE4 (results not shown). The 4.5 kb transcript may result from either alternative mRNA splicing, alternate promoter usage or use of a different polyadenylation motif. For confirmation of the Northern blots, RT-PCR was performed with the PC8-specific primers oPC33.
and oPC35 under saturating PCR conditions, and resultant products were verified by hybridization analysis with an internal oligonucleotide, oPC34 (Figure 4B). All tissues were found to express PC8 mRNA. The ability of the primer pairs to specifically amplify PC8 mRNA from human and rat cDNA suggests that the sequences are well conserved between the two species; the primers did not amplify cDNA for PC1, PC2, PACE4 or furin from either species (results not shown). Since PC8 is widely expressed in endocrine and non-endocrine tissues, it is likely that PC8 is an enzyme of the constitutive pathway of protein secretion and is unlike PC1 and PC2 which are involved in the regulated pathway and have a distribution restricted to endocrine and neuroendocrine cells.

From a comparative perspective, PC8 shares more features in common with furin and PACE4 than with PC1 and PC2. These include: (i) high amino acid identity over the entire molecule with furin and PACE4, whilst the similarity to PC1 and PC2 is lower; (ii) like furin and PACE4, PC8 contains a long (42-amino-acid) signal peptide, whereas the signal sequences of PC1 and PC2 are relatively short; (iii) in common with furin, PC8 possesses a putative transmembrane domain, suggesting that it may be localized to the Golgi; and (iv) the mRNA for PC8 is ubiquitously expressed like those for furin and PACE4, which is in marked contrast with the mRNAs for PC1 and PC2, which are restricted to endocrine and neuroendocrine cells. Combined, these data suggest that PC8 is more closely related to furin and PACE4 than to PC1 and PC2. As such, PC8, furin and PACE4 may have been derived from a common ancestral gene which has been duplicated on chromosome 15 for furin and PACE4 and on chromosome 11 for PC8. It remains to be established whether PC8 can fulfill the functions of furin, whether it does indeed act as a constitutive pathway enzyme and whether it has a substrate sequence specificity for cleavage of proteins similar to that of furin or other members of the convertase family.

This work was supported by a Program Grant from the National Health and Medical Research Council of Australia, by a grant from Chugai Pharmaceutical Co., Ltd., Japan, to T. J. M., J. M. M. and M. T. G., and by Grant GREG 43 to P. B. We are grateful to Dr. D. Stapleton for his helpful discussions.

REFERENCES


Figure 3 Complete amino acid sequence predicted for PC8 aligned to furin, PACE4, PC1 and PC2

The subtilisin-like catalytic domains are boxed and the catalytically important Asp, His, Asn (Asp), and Ser residues are indicated by bold with gaps introduced to maximize the alignments. Residue numbers are indicated at right.

Received 11 December 1995/11 January 1996; accepted 11 January 1996