

Full Length Research Paper

Prediction and analysis of the secretome in *Corynebacterium glutamicum* ATCC 13032

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Corynebacterium glutamicum is an outstanding organism used for amino acid production. Its ability to secrete L-glutamate has been known for almost fifty years now. The complete nucleotide sequence of *C. glutamicum* ATCC 13032 genome was previously determined and allowed the reliable prediction of 3056 protein-coding genes within this genome using computational methods. The 3056 open reading frames (ORFs) of *C. glutamicum* ATCC 13032 were used for the prediction of secreted proteins by bioinformatics approaches, such as SignalP 3.0 and Proteome Analyst. 167 proteins were predicted to be secreted and contain signal peptides, whose amino residues were relatively conserved. Among them, 10 have RR-motif signal peptide and 46 have SignalPasell signal peptide. Total of 167 secreted proteins have functional descriptions, many of which were enzymes that are involved in metabolism. This prediction method has given good insights into the whole secreted proteome of *C. glutamicum* and provided basis to further studies of its secretome features at a genome level.

Key words: *Corynebacterium glutamicum*, secreted proteins, signal peptides.

INTRODUCTION

Corynebacterium glutamicum is a Gram-positive, non-sporulating bacterium that was isolated by Kinoshita and co-workers in a screen for bacteria that secrete L-glutamate. It is used for the industrial production of amino acids such as glutamate and lysine that have been used in human food, animal feed and pharmaceutical products for several decades. In addition, recent studies have indicated the potential of *C. glutamicum* for production of other commercially relevant compounds, such as succinate or ethanol (Bott, 2007; Leuchtenberger, 1996).

Because of the importance of *C. glutamicum* in industrial biotechnology, its 3.3-Mb genome sequence has been determined several times independently. The establishment of a completely annotated *C. glutamicum* genome sequence is a big leap forward to the understanding of the biology of this organism (Kalinowski et al., 2003). The complete genome sequence is the basis for

extensive expression analyses by proteome and transcriptome technologies, which will lead to a comprehensive systemic understanding of gene expression and regulatory networks (Becker et al., 2007).

It is well established that *C. glutamicum* can secrete certain proteins to high concentrations in the medium. However, until recently it was difficult to estimate the number of exported proteins belonging to the secretome of *C. glutamicum*. The completion of the *C. glutamicum* genome sequencing project and the availability of programs for the identification of signal peptides and trans-membrane segments in large collections of protein sequences through worldwide web servers have now made it possible to predict the most likely location of all 3,056 annotated proteins (that is, the proteome) of this organism. Computer-assisted studies have indicated that approximately 15% of the proteome of a given organism, such as *C. glutamicum*, contains membrane sorting signals in the form of hydrophobic stretches of amino acids that can integrate in and span the membrane (Saleh et al., 2001). Some of these putative membrane proteins contain amino-terminal signal peptides (SPs) and may in

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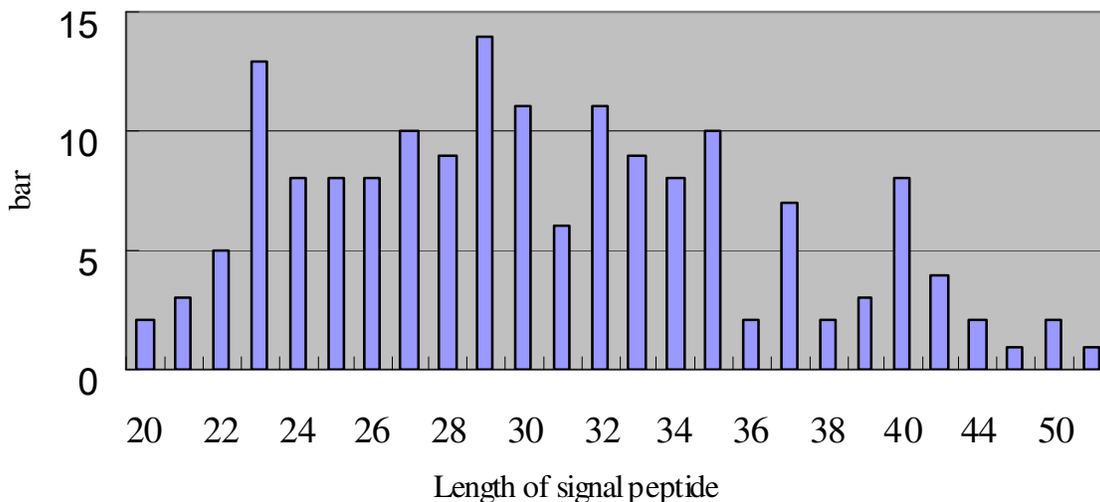


Figure 1. Distribution of signal peptide with different length.

fact be exported proteins.

MATERIALS AND METHODS

Materials

The genome and proteome sequence of *C. glutamicum* ATCC 13032 (accession no. NC_006958) was retrieved from National Center for Biotechnology Information website (<http://www.ncbi.nlm.nih.gov>). Signal peptides and secreted proteins searches were carried out using SignalP 3.0 (<http://www.cbs.dtu.dk/services/SignalP>) and Proteome Analyst (<http://www.cs.ualberta.ca/~bioinfo/PA>), respectively. Lipoprotein SPs and twin-arginine translocation SPs were made with the LipoP 1.0 (<http://www.cbs.dtu.dk/services/LipoP>) and TatP 1.0 (<http://www.cbs.dtu.dk/services/TatP>).

Prediction of SPs using SignalP 3.0 (Bendtsen et al., 2004; Emanuelsson et al., 2000)

SignalP 3.0 server predicts the presence and location of signal peptide cleavage sites in amino acid sequences from different organisms: Gram-positive prokaryotes, Gram-negative prokaryotes and eukaryotes. The method incorporates a prediction of cleavage sites and a signal peptide/non-signal peptide prediction based on a combination of several artificial neural networks and hidden Markov models. Because of the length restrictions, we divided the sequence into 6 submissions to classify proteins as asignal peptides/non-signal peptides.

Prediction of secreted proteins using proteome analyst 2.5

Proteome analyst is a publicly-available, high-throughput, Web-based system for predicting various properties of each protein in an entire proteome (Szafron et al., 2004). We uploaded a FASTA format file containing the 454 SPs sequences to be classified by SignalP 3.0. With the removal of the transmembrane proteins, a set of 167 secreted proteins were obtained and functionally classified. Motif search of secreted proteins uncharacterized was carried out using MYHITS (http://myhits.isb-sib.ch/cgi-bin/motif_scan).

Prediction of lipoprotein signal peptides using LipoP 1.0

The hidden Markov model (HMM) was able to distinguish between lipoproteins (SPaseII-cleaved proteins), SPaseI-cleaved proteins, cytoplasmic proteins and transmembrane proteins. The HMM was able to identify 92.9% of the lipoproteins included in a Gram-positive test set. The results obtained were significantly better than those of previously developed methods (Juncker et al., 2003).

Prediction of twin-arginine signal peptides using Tap 1.0

The method is able to discriminate Tat signal peptides from cytoplasmic proteins carrying a similar motif, as well as from Sec signal peptides, with high accuracy (Nielsen et al., 1999).

RESULTS

Length distribution of predicted signal peptides

The 167 predicted signal peptides (Figure 1) had a length varying from 20 to 55 residues, with an average of 31 residues.

The frequency of 20 amino acids residues of signal peptides

The frequency of 20 amino acids residues of predicted signal peptides is shown in Figure 2. It has to be noted that nonpolar amino acid residues can be found more abundant, where alanine (Ala) residue is most abundant (18%). The frequency of positively charged residues arginine (Arg), lysine (Lys) and histidine (His) is respectively 5.0, 3.7 and 0.8%, while that of negatively charged residues asparagine (Asp) and glutamine (Glu) is 1.2 and 1.6% respectively. For uncharged residues, the frequency of serine (Ser) is the most (9.7%) while tyrosine (Tyr) is

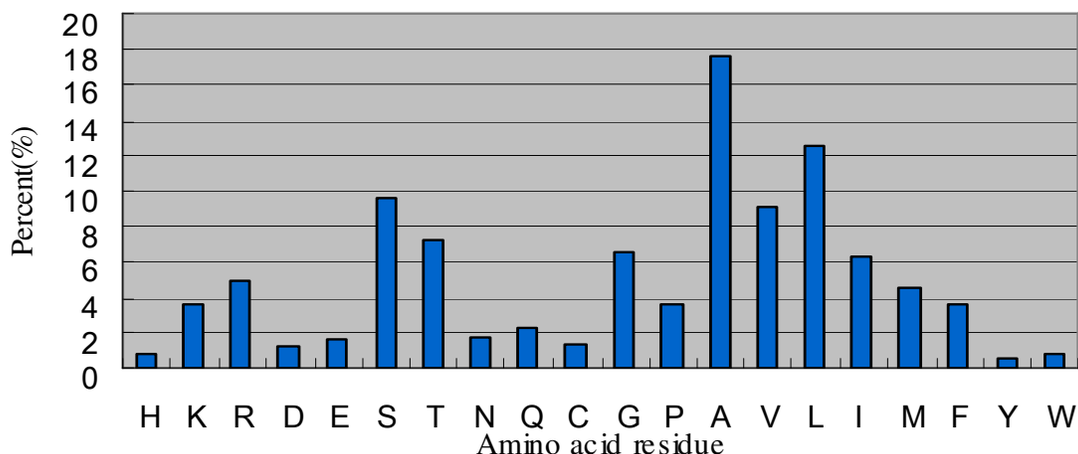


Figure 2. The frequency of 20 amino acids residues of signal peptides in secreted proteins.

Table 1. The frequency of 20 amino acid residues around the signal peptidase cleavage sites (%).

| Amino acid | First letter abbreviation | -3 | -2 | -1 | 1 | 2 |
|---------------|---------------------------|----------|----------|----------|----------|----------|
| Histidine | H | 0 | 1.796407 | 0 | 1.796407 | 0 |
| Lysine | K | 0 | 2.39521 | 0 | 1.197605 | 0 |
| Arginine | R | 0 | 1.197605 | 0.598802 | 1.197605 | 1.197605 |
| Aspartic acid | D | 1.197605 | 1.197605 | 2.994012 | 8.383234 | 8.982036 |
| Glutamic acid | E | 2.39521 | 4.790419 | 0.598802 | 9.580838 | 13.17365 |
| Serine | S | 7.185629 | 17.36527 | 4.191617 | 10.77844 | 16.16766 |
| Threonine | T | 3.592814 | 3.592814 | 0 | 5.389222 | 8.982036 |
| Asparagine | N | 0.598802 | 5.389222 | 1.796407 | 1.197605 | 2.994012 |
| Glutamine | Q | 1.796407 | 10.17964 | 1.796407 | 16.16766 | 1.796407 |
| Cysteine | C | 0.598802 | 5.988024 | 1.197605 | 2.39521 | 4.790419 |
| Glycine | G | 5.988024 | 5.389222 | 5.389222 | 4.191617 | 4.191617 |
| Proline | P | 1.197605 | 2.39521 | 2.39521 | 2.39521 | 12.57485 |
| Alanine | A | 51.49701 | 7.185629 | 74.8503 | 20.95808 | 4.790419 |
| Valine | V | 14.97006 | 8.982036 | 0.598802 | 3.592814 | 8.383234 |
| Leucine | L | 5.389222 | 7.185629 | 0.598802 | 4.790419 | 2.39521 |
| Isoleucine | I | 2.39521 | 6.586826 | 1.197605 | 1.796407 | 2.994012 |
| Methionine | M | 0.598802 | 1.197605 | 0 | 0.598802 | 1.197605 |
| Phenylalanine | F | 0 | 2.39521 | 0.598802 | 2.994012 | 1.796407 |
| Tyrosine | Y | 0 | 1.197605 | 0.598802 | 0.598802 | 1.796407 |
| Tryptophan | W | 0.598802 | 3.592814 | 0.598802 | 0 | 1.796407 |

the least (0.5%). The fact that the residues whose frequency is more than 5% are mostly aliphatic amino acids suggests that such residues are involved in the targeting of secreted protein to specific membrane locations in *C. glutamicum*.

Change of amino-acid residues in C-domain

Three distinct regions comprise the N-terminal signal sequence: the charged N-terminus (N-domain), the

hydrophobic core (H-domain), and the C-terminal cleavage domain (C-domain) (M Akita, 1990; Paetzel et al., 1998). The C-domain of the predicted signal peptides carries a type I SPase cleavage site, with the consensus sequence A-X-A at position -3 to -1 relative to the SPase I cleavage site (Table 1). The frequency of Ala at -3, -1, +1 position is respectively, 51.5, 74.9 and 21.0%. It is important to note that the C-domain must have an extended (β -sheeted) structure for effective interaction with the active site of type I SPases. Based on the crystal structure of the type I SPase of *Escherichia coli*, the side chains of

Table 2. RR-motif signal peptide.

| ID of protein | Signal peptide |
|---------------|------------------------------------------------------|
| gi_62390270 | MAQISRRHFLAAATVAGAGATLAA CA |
| gi_62390439 | MLPIWMGLPFFKAGALSRRKAVFSALGAAALIGAALPTIPTAQA QT |
| gi_62390553 | MVSRRGFLGGAGLIAGASALA GC |
| gi_62391106 | MPQLSRRQFLQTTAVTAGLATFAGTPARA EE |
| gi_62391279 | MFKKHRHGLGSPETKPRSITRRFFTTAAAATLAGLAVLSGCTAQPSQA ED |
| gi_62391337 | MLNIARNRNMKRRLAIAAFVATATATATMAPASA QT |
| gi_62391490 | MSTTITRRNFLRATGILGVAAGIGATLAACA PD |
| gi_62391492 | MRRKLTTLLENKPGARLGGFRALAPTSKIALVLLLLIFLLAIFAPLIKY DP |
| gi_62391811 | MTSSFRRQFLGGLVLAGTGAVA AC |
| gi_62391935 | MVRSTGSMIAITLLSRITGFLRTVMIGAALSPAISA FN |

residues at the -1 and -3 positions are thought to be bound in two shallow hydrophobic substrate-binding pockets (S1 and S3) of the active site, whereas the side chain of the residue at position -2 is pointing outwards from the enzyme. It is presumably for this reason that residues tolerated at positions -3 and -1 of the signal peptide are generally small and uncharged, while almost all residues seem to be allowed at position -2. Nevertheless, a preference for Ser (17%) at position -2 of the signal peptide seems to exist in *C. glutamicum*.

According to the predictions, an Ala residue is most abundant (21%) at position +1 of the mature protein, but all other residues, with the exception of tryptophan (Trp), seem to be allowed at this position.

Lipoprotein signal peptides

Putative lipoprotein signal peptides were identified through similarity searches in the LipoP 1.0 database. Putative lipoprotein sorting signals identified by the method were combined with those identified by SignalP, resulting in a total number of 46. Signal peptides from lipoproteins differ in several respects from those of secretory signals.

Twin-arginine signal peptides

Proteins containing a signal peptide with the RR-motif (R-R-X-#-#, where # is a hydrophobic residue) may be transported via the Tat pathway. Through TatP 1.0 database search for the presence of this motif in amino-terminal protein sequences, a total number of 10 putative RR-signal peptides were identified (Table 2). Notably, the RR-motif was also found in the signal peptides of two putative lipoproteins, suggesting that these proteins might also be substrates for the Tat pathway.

Functional classification of predicted secreted proteins

A total of 167 secreted proteins have functional descrip-

tions, of which 51 are secreted proteins, while 35 are transport system proteins, 43 are enzymes, 18 are binding proteins and precursors (Figure 3).

In order to identify potential functions of 51 secreted proteins uncharacterized, motif analysis was carried out using MYHITS. As a result, five sequences exist as related activity site (Table 3).

Secretion by *C. glutamicum* of heterologous proteins

C. glutamicum has been used for the industrial production of amino acids for several decades. However, there had been only few reports concerning heterologous protein secretion in *C. glutamicum*. Recently, in many studies it has been demonstrated that *Streptomyces mobaraensis* transglutaminase (Date et al., 2004; Kikuchi et al., 2003) another enzyme used in the food industry, and human epidermal growth factor can be efficiently secreted in active form by *C. glutamicum*; so this strain is a potential host for industrial-scale protein production. In addition, the Tat pathway in *C. glutamicum* has been demonstrated to specifically mediate the secretion of *Arthrobacter globiformis* isomaltodextranase and green fluorescent protein (GFP) carrying an *E. coli* TorA signal peptide. Furthermore, the Tat-pathway-dependent secretion of GFP has been shown to be far superior in *C. glutamicum* compared with two other Gram-positive bacteria, *Bacillus subtilis* and *Staphylococcus carnosus* (Meissner et al., 2007). More recently, there was report of secretion of *Streptococcus bovis* α -amylase using cspB promoter and signal sequence by *C. glutamicum* for the efficient utilization of raw starch, identifying *C. glutamicum* as a very useful host for the expression of heterologous proteins (Kikuchi et al., 2006; Tateno and Akihiko, 2007; Kikuchi et al., 2007).

In the present prediction, about 21% of secreted proteins were involved in transport system, while 10 putative RR-signal peptides were identified. Thus, the observations that *C. glutamicum* can efficiently release different heterologous proteins into the culture medium subsequently to their Tat-dependent translocation across

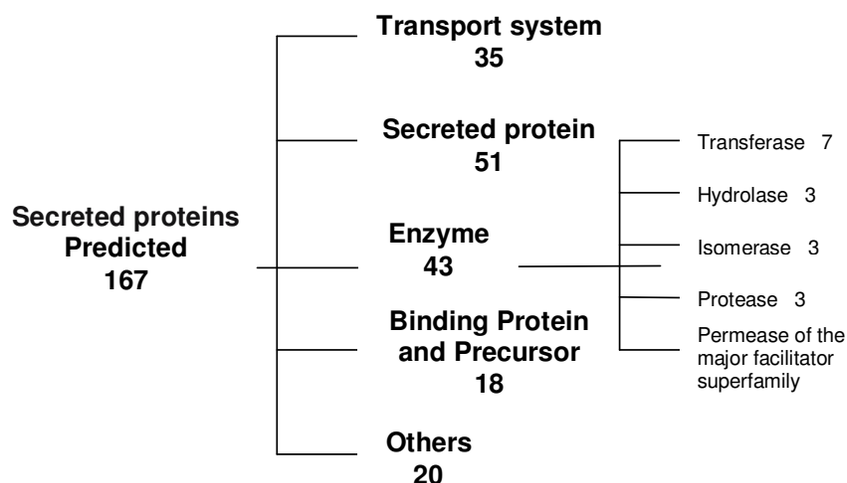


Figure 3. Classification of predicted secreted proteins.

Table 3. Sequences with key motifs in secreted proteins with unknown function.

| ID of protein | Description of the motif | Possible function |
|---------------|----------------------------------------------------------|------------------------------------------------------|
| gi_62389680 | Serine proteases, trypsin family, histidine active site. | Serine protease |
| gi_62390357 | Hemopexin domain signature. | Hemopexin |
| gi_62391502 | TonB-dependent receptor proteins signature 1. | TonB-dependent receptor |
| gi_62391027 | Transcription factor TFIIB repeat signature. | a target of gene-specific transcriptional activators |
| gi_62391174 | Aldehyde dehydrogenases glutamic acid active site. | Aldehyde dehydrogenase |

the plasma membrane is somewhat unexpected, particularly in view of the fact that the proteins translocated via the Tat pathway usually arrive at the transside of the cytoplasmic membrane in a fully folded state.

Identified enzymes in the predicted secreted proteins

Extensive works have been done to investigate other functions of *C. glutamicum* besides amino acids production binding with the enzyme protein secreted, such as aromatic degradation. In their work, many secreted proteins were identified (Table 4), while some of which were also identified by our prediction.

DISCUSSION

In the present work, efforts were made to identify all genes in the *C. glutamicum* genome database whose deduced proteins would likely be soluble secreted proteins (the secretome). While certain *C. glutamicum* secretory proteins have been studied in detail, such as six mycolyltransferase genes and their gene products, more data on the entire secretome is needed. One approach to rapidly predict the functions of an entire proteome is to utilize genomic database information and

prediction algorithms. The use of computer-based prediction algorithms is a powerful, systematic, and rapid tool to obtain preliminary functional information on gene products of an entire genome. Information can then be analyzed in global fashion to organize functional groupings of predicted proteins, or individually, in order to identify genes of particular interest for future experimental study. The *C. glutamicum* genome database was queried in an effort to identify all genes whose deduced proteins would likely be secreted proteins in order to: (a) obtain a global perspective on secreted proteins in *C. glutamicum*; and (b) identify previously uncharacterized genes for further experimental study. A series of prediction algorithms available was therefore used on internet-based servers to analyze the *C. glutamicum* genome database.

In this study, identification was carried out on genes whose proteins have signal peptides and are known to be secreted extracellularly, including: *cop1*, *cmt1*, *cmt2*, *cmt3*, *cmt4*, and *cmt5*. Interestingly, 42 of secreted proteins predicted are of unknown function. In order to gain additional insight into the functional properties of these potential *C. glutamicum* secretory proteins in our dataset, we referred to the extensive motif search and got some related activity site, such as serine protease, Aldehyde dehydrogenase.

Important limitations of this approach are that it relies on prediction algorithms with a defined error rate which

Table 4. Identified enzymes in the predicted secreted proteins.

| ID of protein | Enzymes | EC Number | Reference |
|---------------|----------------------------------------------------------|-----------|----------------------------|
| gi_62389241 | Trehalose corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62389812 | corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62389917 | corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62391021 | corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62391714 | Trehalose corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62391716 | Trehalose corynomycolyl transferase | 2.3.1.122 | (Brand et al., 2003) |
| gi_62391453 | putative secreted protein, hypothetical endoglucanase | | (Brand et al., 2003) |
| gi_62390420 | secreted cell wall-associated hydrolase | | (Brand et al., 2003) |
| gi_62389427 | secreted protein | | (Brand et al., 2003) |
| gi_62390276 | putative secreted hydrolase | | (Brand et al., 2003) |
| gi_62389528 | ABC-type amino acid transport system, secreted component | | (Schluesener et al., 2007) |
| gi_62390348 | NADH Dehydrogenase | 1.6.99.3 | (Schluesener et al., 2007) |
| gi_62390845 | putative secreted or membrane protein | | (Schluesener et al., 2007) |
| gi_62389441 | Probable short-chain dehydrogenase, secreted | | (Huang et al., 2008) |

could potentially be greater in specific organisms. Furthermore, these prediction algorithms are useful for rapid preliminary analyses of large amounts of genomic data, but it must be emphasized that these are only predictions, which require experimental validation. The present approach was to be inclusive rather than exclusive; so overall, these results probably represent an overestimation of the actual *C. glutamicum* secretome, especially since many open reading frames (ORFs) in the genome database have not been confirmed experimentally and some ORFs may not be expressed.

In conclusion, for further experimental study, we would like to examine novel secreted proteins and identify their function using proteomics-based approaches to analyze *C. glutamicum* secreted proteins.

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