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Elucidating the Role of CaM KK in Cell Cycle and Cell Fate Using a C. elegans Model

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The mammalian calmodulin-dependent protein kinase kinases (CaM KKs) have been shown to phosphorylate and activate the calmodulin-dependent protein kinases I and IV (CaMKI, CaMKIV), leading to transcription of a variety of genes important in cell cycle regulation. Using a C. elegans homologue (ceCaM KK), we are investigating the importance of this proposed cascade to cell cycle, cell fate and development in the context of a well-defined multicellular organism. By generating transgenic worms with reporter proteins controlled by the ceCaM KK promoter, gene expression has been demonstrated in the excretory cell, vulval muscle cells and several neurons of adult hermaphrodites, in the hypodermal cells of L1/L2 larvae, and in several male-specific tail cells. To examine this pathway biochemically, we have cloned the ceCaM KK and ceCaMKI cDNAs, and produced recombinant proteins by prokaryotic expression methods. Both mammalian and C. elegans CaM KKs can phosphorylate either species' CaMKI homolog specifically on the activation loop (T177 in human, T179 in C. elegans) in vitro. These results indicate a functional homology between the mammalian and C. elegans calmodulin dependent kinases, and demonstrate the existence of a calmodulin dependent kinase cascade in the worm. Further work remains to reveal the biological functions of this signaling.
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Annual Report for Grant DAMD17-97-1-7331
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Ethan E. Corcoran

Table of Contents

Introduction 2
Results and Discussion 2
Conclusions 5
Appendix: Key Accomplishments 6
Reportable Outcomes 6
Acronyms/Abbreviations 6
References 7
Introduction

Calcium and calmodulin (CaM) are ubiquitous signaling molecules, implicated in a wide variety of cellular functions including cell cycle control. Ca\(^{2+}\)/CaM signals are critical in G\(_0\) reentry, G\(_1\)/S, G\(_2\)/M, and metaphase transitions in a variety of organisms (1). Recently, a novel kinase cascade has been proposed connecting the action of CaM dependent kinase kinases (CaMKK) to transcription of key factors in cell cycle control, through the phosphorylation and activation of calmodulin dependent kinases I and IV (CaMKI and CaMKIV) (2-7). In addition, mammalian CaMKK has now been implicated in the regulation of protein kinase B and of the mitogen-activated protein kinase (MAPK) cascade, both of which have significant established roles in cell cycle control and apoptosis (8-9). This project was designed to assess the biological relevance of these relationships and of the CaMKK itself. Using a Caenorhabditis elegans model, any perturbation of normal cell division or cell fates can be identified (10-12). We have now cloned cDNAs for C. elegans isoforms of both CaMKK and CaMKI (ceCaMKK and ceCaMKI), examined their biochemical homology using recombinantly expressed proteins, and used transgenic techniques to determine the developmental expression pattern of ceCaMKK. We are currently pursuing the downstream effectors of this cascade through an in vitro screen for kinase substrates of either protein, and attempting to isolate null mutants for these genes. Understanding this kinase cascade, from protein interactions to biological consequences, may help unravel the essential functions of calcium signals in the cell cycle.

Results and Discussion

As indicated in last year’s report, the first project objective of cloning ceCaMKK has been completed. While we have not empirically confirmed the sequence of the last 80 bp of the 5’ end, a recent report successfully used rtPCR to clone the predicted C05H8.1 gene (13). Unfortunately, while this study does verify the 5’ start sequence, the 3’ primers included an assumed TAA stop codon. This oversight led to the conclusion that the transcript is identical to the predicted open reading frame, and missed the final two exons of the gene. Since our cloning strategy involved hybridization screening of a cDNA library, and did not assume the location of the stop codon, we have avoided this pitfall (14). Interestingly, the only biochemical difference demonstrated so far between the full length ceCaMKK and this truncated C05H8.1 is autophosphorylation of the truncated form (13).

The second objective in this project, determining expression patterns and isolating null mutants, has presented the greatest difficulties to date. In last year’s report, the developmentally regulated expression pattern for hermaphrodite worms had been examined using transgenic methods (15). Expression is observed in the adult vulval muscle cells, amphid sensory neurons and the excretory cell, as well as in L1 hypodermal cells. Using the same transgenic constructs, the male nematodes’ expression pattern has been shown to include the excretory cell, amphid sensory neurons, and several male specific tail cells, believed but not yet confirmed to be neurons. Larval males have not been examined. To extend these results, transgenic constructs that include the entire ceCaMKK gene (promoter and coding regions) in fusion with GFP are being made. These constructs will allow the subcellular localization of the protein to be determined, as a function of development and stimulation of the relevant neurons in adults.
The difficulty with these methods for determining expression has been an inability to confirm the patterns using a second technique. In-situ hybridization has been attempted using standard methods and three different gene specific DNA and RNA probes without success, most likely because the expression levels of the ceCaMKK are so low (16-17). Our labs are able to use these methods for more strongly expressed signals, such as RNAs induced by cadmium exposure (18). No specific antibody is yet available, so immunohistochemical methods have not been tried. The increasingly frequent use and confirmation of transgenic methods for studying expression patterns makes secondary confirmation of this pattern less important than it was when the project was first proposed. However, being unable to confirm the presence or absence of either RNAi message or protein in a worm makes it difficult to determine the success of attempts to use RNAi suppression of the gene.

The second focus of objective two, obtaining and characterizing null mutants of ceCaMKK, is still in progress. Two methods are being employed - PCR screening of mutagenized worm libraries and RNAi. PCR screening of mutant libraries requires the construction of a mutagenized deletion worm library, which can be generated using well-established techniques (19). Once the worms have been mutagenized, they are split into small groups, grown for one generation, and divided into a freezer stock and a DNA stock. The DNA stock is used to make genomic templates for PCR reactions, while the freezer stock is saved for the recovery of any relevant worm mutants. A two step nested PCR strategy is used to identify any deletions within the region of interest, which can be followed until a unique strain has been isolated from the mixture. Our lab participated in the construction of such a library in collaboration with four other labs at Duke. PCR screening of this library has not yet yielded any relevant mutants. The same basic strategy has been adopted by the Sanger Center’s C. elegans Gene Knockout Program, with the goal of eventually obtaining knockouts of every known worm gene (20). We have requested that they screen their libraries for deletions of the ceCaMKK gene, but, at last report, two libraries had been screened with no positives. The Sanger Center is currently working on methods to improve the library generation and screening procedures, and will continue to search for deletions in the ceCaMKK locus.

The alternate method for obtaining null mutants in a gene involves the poorly understood phenomenon of RNAi. It has been shown for many early larval genes that injection of double stranded RNA effectively abolishes the expression of that gene in the progeny of injected worms (21-23). The effectiveness of the expression block varies with the gene, the dsRNA used, the cells involved and the stage of development. It is most effective during early development, but has been observed to cause adult phenotypes and sometimes persist into another generation (21). We have generated and injected dsRNAs corresponding to the ceCaMKK mRNA, but no phenotype has been observed. This could indicate that there is no developmental consequence of ceCaMKK gene suppression, that the consequence is not obvious without appropriate testing, or that the gene has not been successfully suppressed by the dsRNA. As mentioned previously, we are continuing to work on a method for measuring expression levels in situ in order to determine the success or failure of RNAi in this case. The search for null mutant of the ceCaMKK will continue to be a major focus of the project during the next year.

The final objective of the project is to identify other components of the ceCaMKK pathway. As mentioned last year, by BLAST searching the C. elegans genomic sequence, a potential homologue of CaMKI was identified in the predicted gene K07A9.2 (24). The open reading frame appeared to be missing the 5’ end, including the ATP binding loop. This may have been due to uncertainty in the genomic sequence 5’ to the gene. Initially, the predicted
gene was cloned by rtPCR, and 5' RACE was used to find the remaining 5' region. The 5' RACE products were all short sequences, ending at similar sites only 20 bases upstream of the predicted start. Searching upstream genomic sequences for homology to the 5' end of CaMKI revealed two additional potential exons beginning more than 8 kb 5' of the predicted K07A9.2 start site. By rtPCR, these exons were amplified as part of a complete ceCaMKI mRNA, having 58% amino acid identity to human CaMKI (30% to human CaMKIVa). Nonetheless, the shorter transcript, corresponding to the predicted K07A9.2 gene, includes sequences not found in the longer transcript, has its own start codon, and appears by 5'RACE to have a transcriptional start site. It is possible, therefore, that this gene has two products, one a functional kinase and the other, by truncation of the ATP binding domain, a kinase dead variant. This may be analogous to the mammalian CaMKIV gene, which is known to have multiple products including an active kinase and a truncated form regulated by an internal promoter whose function is unknown (3).

We are working on several methods to examine the ceCaMKI gene, including northern blots using probes to the 3' end of the mRNA and affinity purification of a mammalian CaMKI antibody, which by homology has a good chance of recognizing the worm protein but as a polyclonal antiserum reacts with many proteins in worm extracts. In addition, transgenic expression constructs are being generated which will include various regions of the promoter and first introns, to examine the kinase’s expression pattern and ascertain if there is an internal promoter which might regulate a second gene product.

The full length ceCaMKI cDNA has been used for recombinant expression to study the protein’s biochemistry. It expresses well in bacteria as a GST fusion protein, and can be purified to near homogeneity. When used for kinase assays, it is remarkably similar to human CaMKI. It binds CaM in overlay experiments, is Ca\textsuperscript{2+}/CaM dependent for its kinase activity, and is phosphorylated and activated by either recombinant mammalian CaMKKB or ceCaMKK (25-26). Mutation of threonine 179 to alanine in the activation loop (analogous to T177A in the human form) abolishes phosphorylation by kinase kinases and prevents activation (26). The specific ATP, CaM and peptide substrate binding constants have not yet been determined.

With the identification and expression of the ceCaMKI, the focus for screening the ceCaMKK pathway, objective three, has turned to finding substrates of the kinase cascade. Following the methods successfully applied by Tony Hunter and colleagues in the screen for MAPK substrates (27), we have been developing an in vitro screen for protein substrates of this kinase cascade. We have a unidirectional cDNA library based on the lambda-ZAP-XR vector (Stratagene); this type of library has been used for antibody expression screens. The basic strategy involves inducing a growing phage cDNA library with IPTG impregnated nitrocellulose membranes. After 4-6 hours at 30 degrees, these filters are removed and a duplicate filter is placed on each plate for a further 4 hours. The filters bind proteins from the bacterial lysates made by the lytic phage infection, including the expressed cDNA specific to each phage. The filters are blocked, washed and incubated with ATP (to avoid isolating ATP-binding proteins), before they are exposed to activated kinase in the presence of \(\gamma^{32}\text{P}-\text{ATP}\). After further washes to remove unincorporated isotope, the filters are exposed to film, and positives appearing on both duplicate filters can be selected for further screening. The reaction conditions have been adapted to accommodate the calmodulin-dependent protein kinases by including calcium, calmodulin and increased ATP in the reaction buffer and by preincubating the kinase with kinase kinase at high ATP concentrations to activate the kinase. The ceCaMKK has been difficult to purify and is only produced in small quantities, so recombinant mammalian CaMKKB is used to activate
ceCaMKI used for the screen. Samples of the final kinase buffer are very reactive against purified substrates.

To test these methods, a lambda-ZAP (Stratagene) control phage containing a convenient protein substrate sequence as an insert (the 1-117 amino acid fragment of p300, referred to as \( \lambda 117 \)) (28). Bacteria infected with \( \lambda 117 \) have been used in comparison with bacteria infected with wild type phage (\( \lambda \text{WT} \)) to test conditions for the screen. At this point, \( \lambda 117 \) and \( \lambda \text{WT} \) derived bacterial extracts are easily identified on a gel assay, with a very strong band corresponding to the 1-177 p300 fragment, but when immobilized on filters, the signal from \( \lambda 117 \) is difficult to distinguish from \( \lambda \text{WT} \). It should be possible to adjust conditions so that the background is sufficiently distinguishable from a positive signal. Using these reagents, screening conditions will be optimized until the signal to background ratio is high enough to permit a full scale screen of the cDNA expression library. Since we are screening a \( C. \text{elegans} \) library, not mammalian, fewer plaques need to be screened for complete coverage of the genome, and any substrate cDNA fragments identified should be more easily cloned using information from the cDNA and genome sequencing projects.

Conclusions

We have identified and cloned a calmodulin dependent protein kinase cascade in \( C. \text{elegans} \). Using recombinant expression, we have confirmed biochemical homology between the \( C. \text{elegans} \) and mammalian CaMKK and CaMKI, and are continuing to search for targets of the cascade by an in vitro screening method. To define biological functions of this cascade, we are investigating the expression patterns and gene regulation of these kinases and attempting to generate null mutants of these genes using the available reverse genetic techniques. Completion of this project will be a significant advance in our understanding of the biological roles of calcium, calmodulin and calmodulin dependent protein kinases.
Appendices

Key Research Accomplishments

1. Cloning of the \textit{C. elegans} homologue of the calmodulin dependent protein kinase kinase (ceCaMKK)
2. Prokaryotic expression of recombinant ceCaMKK
3. Confirmation of biochemical homology between mammalian CaMKK and ceCaMKK
4. Determination of the developmental and cell specific expression pattern of ceCaMKK in \textit{C. elegans} using transgenic methods
5. Cloning of the \textit{C. elegans} homologue of the calmodulin dependent protein kinase I, ceCaMKI
6. Prokaryotic expression of recombinant ceCaMKI as a GST fusion protein
7. Confirmation of biochemical homology between mammalian CaMKI and ceCaMKI
8. Development of an in vitro screening protocol for finding substrates of the CaM dependent kinases

Reportable Outcomes

None to date.

Acronyms/Abbreviations

\begin{itemize}
  \item ATP \hspace{1cm} \text{adenosine triphosphate}
  \item \textit{C. elegans} \hspace{1cm} \textit{Caenorhabditis elegans}
  \item CaM \hspace{1cm} \text{calmodulin}
  \item CaMKI \hspace{1cm} \text{calmodulin dependent kinase I}
  \item CaMKIV \hspace{1cm} \text{calmodulin dependent kinase IV}
  \item CaMKKB \hspace{1cm} \text{calmodulin dependent kinase kinase B}
  \item ceCaMKK \hspace{1cm} \textit{C. elegans} calmodulin dependent protein kinase kinase
  \item ceCaMKI \hspace{1cm} \textit{C. elegans} calmodulin dependent protein kinase I
  \item CREB \hspace{1cm} \text{cyclic adenosine 3',5'-monophosphate response element binding protein}
  \item GFP \hspace{1cm} \text{green fluorescent protein}
  \item GST \hspace{1cm} \text{glutathione-S-transferase}
  \item IPTG \hspace{1cm} \text{isopropyl \(\beta\)-D-thiogalactopyronoside}
  \item \(\lambda\)117 \hspace{1cm} \text{lambda-ZAP phage expressing the 1-117 amino acid fragment of p300}
  \item \(\lambda\)WT \hspace{1cm} \text{wild type lambda-ZAP phage}
  \item MAPK \hspace{1cm} \text{mitogen-activated protein kinase}
  \item MBP \hspace{1cm} \text{maltose binding protein}
  \item PCR \hspace{1cm} \text{polymerase chain reaction}
  \item PKB \hspace{1cm} \text{protein kinase B}
  \item RACE \hspace{1cm} \text{rapid amplification of cDNA ends}
  \item RNAi \hspace{1cm} \text{double stranded RNA mediated inhibition (see ref 21)}
\end{itemize}
References


20. A description of this ongoing project is available on the internet at: http://www.sanger.ac.uk/Projects/C_elegans/Knockout/


