

Investigating *KNOX* Gene Expression in *Aquilegia* Petal Spur Development

Imtiyaz Hossain and Clara Levy

Cambridge Rindge and Latin High School, 459 Broadway, Cambridge MA 02138, USA

Kramer Laboratory, Organismic and Evolutionary Biology Department, Harvard University, 16 Divinity Avenue, Cambridge MA 02138, USA

Summary

Organ growth in many flowering plants progresses through two phases: cell proliferation and cell elongation. Recent work has indicated that class 1 *KNOX* genes regulate cell proliferation. These genes encode transcription factors that are largely responsible for a steady supply of undifferentiated stem cells. Moreover, *KNOX* gene expression is detectable in lateral organs of different plants such as compound leaves, suggesting a role in sculpting organ shape. The association of *KNOX* genes with stem cells suggests a role for these genes in the cell division phase of plant development. This suggests that a specific lateral organ, petal spurs, may be built in part using *KNOX* genes. Indeed, previous work has shown that *KNOX* gene over-expression in petals may cause spur-like outgrowths. In *Aquilegia*, a short proliferation phase gives rise to nascent spurs, which is followed by a cell elongation phase. We investigated *KNOX* gene expression and found by using Reverse Transcription Polymerase Chain Reaction, that expression of *KNOX* genes was detectable, but not uniformly so, suggesting that there may be either a highly cell-specific expression of *KNOX* genes or extremely low expression in petal spurs. Future tests with micro dissected tissue samples may prove to be helpful but it can be concluded for now that *KNOX* genes are not highly or consistently expressed in *Aquilegia* petal spurs, and their role in this organ is an ongoing mystery.

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Introduction

Whether or not similar genetic pathways are used to program convergent biological structures poses an interesting evolutionary question. Plants, lacking cellular mobility, are particularly adept at co-opting regulatory pathways and assigning them new roles, resulting in novel structures. For example, the pathway that maintains undifferentiated stem cell fate and then promotes differentiation into lateral organs in the shoot is deeply conserved within plants. At the same time the genetic players have been co-opted several times independently

to function in new spatial or temporal areas, resulting in different patterning of lateral organs (1).

Petal nectar spurs have evolved several times independently across the flowering plants, often times co-evolving with different pollinators (2). The genus *Aquilegia* is a flowering plant that is a useful model system for genetic and genomic studies (3). The petals remain short for a period during development before extending a blade that curves as it elongates, forming a cup-shape that elongates into the spur (4).

All plant shoot tissue is derived from the apical meristem (shown in **Figure 1**), which is comprised of a region of undifferentiated cells that can give rise to any somatic tissue. The cells of this region perpetually divide, providing a steady supply of stem cells, which allows the plant to facilitate growth and produce new organs throughout its lifetime. The genetic machinery that is involved in maintaining meristem indeterminacy or in stopping indeterminacy and promoting differentiation can be turned on in other regions of the plant (5).

In *Arabidopsis*, a close relative, there are 8 *KNOX* genes that are divided into two classes: class I and class II. The class I genes encode four transcription factors: *SHOOTMERISTEMLESS* (*STM*), *BREVIPEDICEL-LUS* (*BP*, also known as *KNAT1*), and *KNOTTED-like* (*KNAT2* and *KNAT6*).

Genes such as *SHOOT MERISTEMLESS* (*STM*) promote cell proliferation in the shoot apical meristem. These genes encode transcription factors, which are proteins that bind to regulatory DNA sequences of downstream targets. This binding allows *KNOX* genes to regulate levels of expression of the downstream target genes.

However, *KNOX* gene function is not limited to the meristems. They have roles promoting compound leaf structure and can partner with another gene family, *ARP* (8). *KNOX* genes must be down-regulated in order for cells in the leaf to differentiate, but if they are reactivated in leaves, this results in different areas of the leaf or leaflets. Studies show that in *Arabidopsis*, expression of class 1 *KNOX* genes in leaves induces formation of compound leaf structure or leaflets (9). Simple leaves showed high levels of *KNOX* gene expression in the meristem and no expression in the site of leaf initiation. After induction of *KNOX* expression, leaves developed with complex leaf phenotypes and various shapes. This shows that there is some relationship between *KNOX* genes and their partners, such as *ARP*, in the development of leaf structure as well as with meristem identity.

If *KNOX* genes can manipulate the growth of these

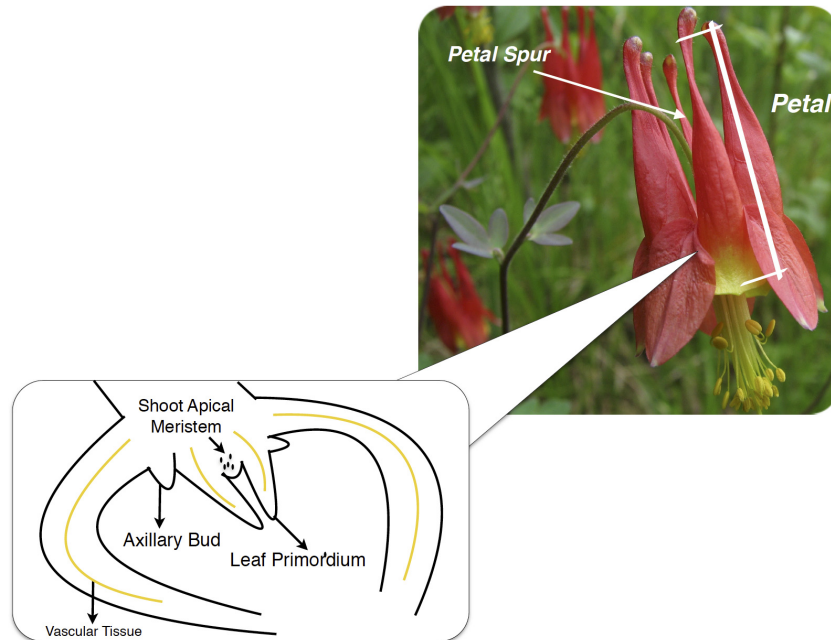


Figure 1: A diagram of *Aquilegia caerulea*. The shoot apical meristem houses many of the undifferentiated cells for the flowering plant. Just along side it are the leaf primordia, which are lateral outgrowths that eventually develop in leaves. Axillary buds grow from the axil of leaf and may develop into another leaf or even a separate flower. When petals fully develop, they tend to create tubular extensions called spurs, which are evolutionarily significant in terms of pollination. This investigation is focused on these spurs and by using RT-PCR, we determined which *KNOX* genes are expressed in this organ. (See 21, 22).

two organs, they may be able to influence the development of other organs, such as petal spurs. Spurs are tubular extensions that grow off of floral organs. They have evolved several times independently across flowering plants. It has been noted that there is a distinct correlation between the highly variable length of spurs in the plant *Aquilegia* and pollinator tongue lengths (10). These outgrowths allow for coevolution with pollinators in the environments, which facilitate rapid speciation among families of flowers (11). For the plant, it would be very efficient to co-opt already present mechanisms for the rapidly evolving organs.

But how would *KNOX* genes play a functional role in the development of spurs? Let's start with how plants generally develop. There are typically two stages of cell growth: cell proliferation and cell elongation (12). The process begins in the apical meristem, in which stem cells divide and give rise to daughter cells. However, if these cells are outside the zone of influence of *KNOX* genes, they are recruited to differentiate into lateral organs. Lateral organs include leaves or floral meristems if the proper environmental cues are present. These floral meristems then give rise to floral organs.

Spur growth could be due to reactivation of *KNOX* transcription factors in petals, which may increase the number of cells in the spur. However, this seems unlikely in *Aquilegia*, as there is evidence of spur development that is void of any *KNOX* gene expression (11). Furthermore, while expression of *KNOX* genes in *Arabidopsis* showed leaf phenotypes analogous to compound leaves, there were no floral phenotypes (13).

Other groups have investigated whether or not the *KNOX* genes that promote stem cell fate in the meristem have been turned on again in petals to prolong cell division and promote extension of the petal into an elongated organ. The role of *KNOX* genes could be different in plants lacking spurs. *Antirrhinum*, better known as snapdragon, does not naturally grow spurs (14). Increased expression of two of its *KNOX* genes resulted in odd spur-like growths that developed off of the petals. Studies executed in *Linaria*, a plant that is closely related to *Antirrhinum*, demonstrated that *KNOX* genes were highly expressed in the floral organs and the front side of the petals known as the "spur-producing ventral petal" (15). Expression of the *Linaria KNOX* genes in tobacco resulted in spur-like outgrowths. However, expression of one of the class 1 *KNOX* genes from the spur producing orchid *Dactylophiza fuschii* in tobacco failed to produce spur outgrowths (15). These studies indicate that *KNOX* genes are expressed in petals of some taxa, and that some of these related genes, but not all, promote outgrowths. If the *KNOX* genes are responsible for spur growth in some taxa, it is not universal. The studies also did not confirm whether the spur like outgrowths were in fact true spurs.

Past research suggests that spur development in *Aquilegia* consists of two phases: an initial phase of cell divisions throughout, and then a second phase of cell elongation (16). The initial phase of cell proliferation cannot account for the entirety of growth of the plant. Cell elongation accounts for the rest of the increase in size of the plant. Other research suggests that the effects

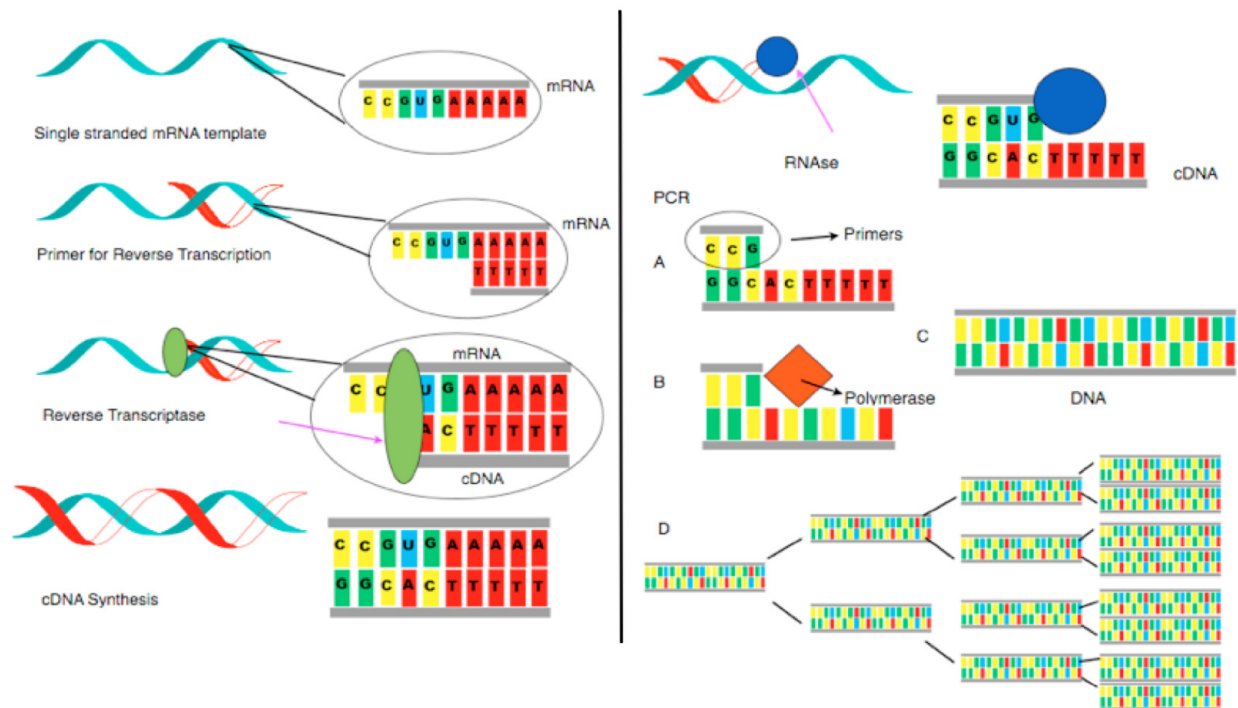
Reverse Transcriptase Polymerase Chain Reaction
RT-PCR

Figure 2: Overview of Reverse Transcriptase Polymerase Chain Reactions (RT-PCR). *Aquilegia* RNA was isolated from 5 organs: sepals, petals, stamens, staminodia, and carpels. In RT-PCR, cDNA is first synthesized using *KNOX* gene primer dilutions, template RNA, and the enzyme reverse transcriptase to add complimentary nucleotide base pairs to the RNA. To initiate PCR, the original RNA template is removed by RNase and cDNA is amplified. PCR then begins, in which annealing occurs with primers, creating a template for Taq polymerase to complete the double stranded cDNA. Denaturation follows, where primers anneal and extend the cDNA for a set number of cycles, creating amplified cDNA. After the reaction is complete, gene expression can be visualized through stained gels.

of *KNOX* genes do not influence petal spur growth in *Aquilegia* (Elena M. Kramer, unpublished data). In fact, they have almost no association with the first phase of petal spur development, cell proliferation (11). These complex lateral organs may just be anomalies in terms of petal spur curvature.

It has also become apparent that cell proliferation is an extremely short phase in *Aquilegia* development (16). A comparison was made to *Linaria*, which have petal spurs as well and results have shown that *KNOX* gene expression indeed exists during its cell proliferation phase. This may suggest that a different set of genes control these spur outgrowths. However, we wanted to expand this study to a second system with spurs. We repeated the gene expression studies done in Collani *et al.* with more petal samples. Confirming these results was an immediate goal because it seemed interesting that the development of similar structures in flowering plants are driven by different mechanisms.

In *Aquilegia*, cell number is irrelevant, and the second phase is responsible for the majority of spur growth. This investigation's purpose is to further our understanding of the role of *KNOX* genes in *Aquilegia*. Since cell divisions are a minor part of spur development, we hypothesized

that *KNOX* genes would not have a role in spur growth. Our results were largely consistent and indicated that the *KNOX* genes likely do not play a significant role in the development of spurs in *Aquilegia* (Elena M. Kramer, unpublished data). However, we found expression of one related gene of a *KNOX* gene in petals.

Results

We chose to use RT-PCR to investigate *KNOX* gene expression in petals because it was a straightforward and fairly quick process (Figure 2) to isolate RNA, synthesize cDNA, and run the PCR. We used five different samples of petals and tested expression of eight different *Aquilegia* genes: *CYCLOIDEA* (*CYC*), *HISTONE4* (*HIS*), *KNOTTED* (*KN*), *KNOTTED-LIKE1* (*KXL1*), *KNOTTED-LIKE2* (*KXL2*), *SHOOTMERISTEMLESS1* (*STM1*), *SHOOTMERISTEMLESS2* (*STM2*), and *TCP4* (Figure 3). For our gene expression analyses, we pooled five groups of petals from different flowers. We tested expression of housekeeping control gene *ISOPENTYL PYROPHOSPHATE: DIMETHYLALLYL PYROPHOSPHATE ISOMERASE2* (*AqIPP2*) as a positive control (Figure 4).

HIS4 encodes a known cell division marker (16).

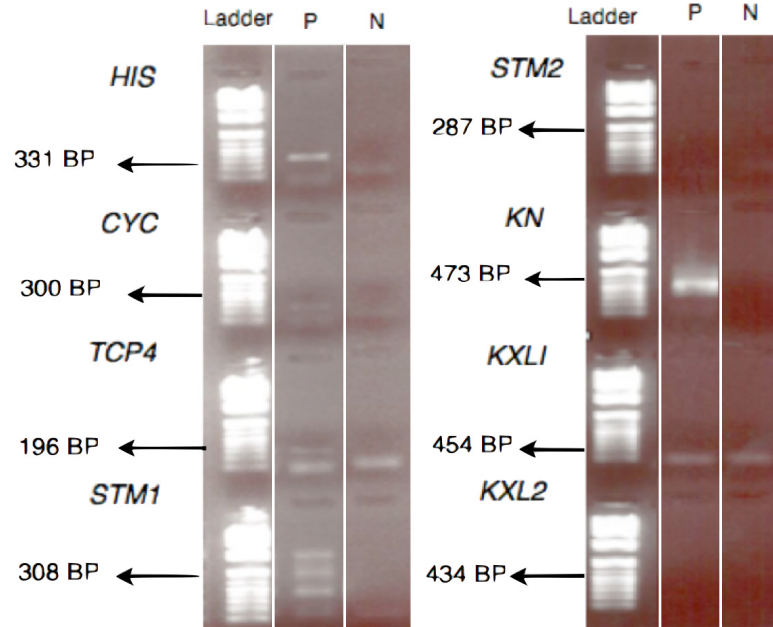


Figure 3: Gene expression in wild type *Aquilegia* petal spurs. After running RT-PCR, gene expression levels of *HIS*, *CYC*, *TCP4*, *STM1*, *STM2*, *KN*, *KXL1*, and *KXL2* in *Aquilegia* petal spurs were shown on a stained gel. Results show consistent expression for *HIS*, *TCP4*, *KN* and *STM1*. However, there is no expression in *CYC*, *STM2* and *KXL2*. P: petal organ. N: negative control (water). BP: base pairs.

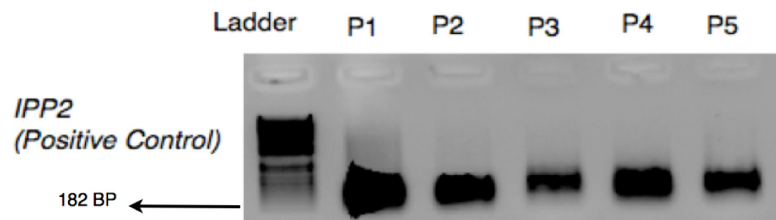


Figure 4: *IPP2* expression in wild type *Aquilegia* petals. Additionally, RT-PCR was performed on *IPP2* and expression levels were shown on a stained gel. *IPP2* is a housekeeping gene for *Aquilegia*, therefore a suitable positive control for the investigation. As expected, *IPP2* was highly expressed in 5 different petal spur samples.

KNOTTED (*KN*) encodes a protein that helps cells perform cytokinesis, a necessary step in cell division. *KXL1* and *KXL2* are *KNOTTED*-like homologs identified in *Aquilegia* (17). *STM1* and *STM2* are the *Aquilegia* orthologs of the *Arabidopsis* *STM* transcription factor that maintains cell indeterminacy in the shoot apical meristem. *CYC* is a transcription factor known to have roles promoting petal lobing and has roles in patterning bilateral symmetry (18). Finally, *TCP4* is a member of the *TCP* gene family (19), which consists of several proteins that share a TCP domain. This transcription factor restricts the domain of cell proliferation (20).

The first lane in our gel image (Figure 3) is a 1 kb ladder that was used to check if our band sizes seen from our RT-PCRs were the size we expected from the primers we designed. The second lane shows our petal RT-PCR products from each experiment done with each primer, and the third lane is a negative control, which was performed using water instead of cDNA.

We found that *HIS* was expressed in petals, which may be an indicator that cell proliferation is active for longer than previously thought. However, *TCP4*, which

acts as an antagonist of cell division, was also moderately expressed. From this we hypothesize that there are different zones of cell division and expansion occurring in our samples. Since we used whole-petal RNA, we hypothesize that *HIS* might still be active in the blade of the petal, while *TCP4* expression is primarily in the spur (Figure 1).

The different *KNOX* genes that we tested and their corresponding levels of expression raise questions on why there is variation between these genes in petal spurs. From our results, *KN* seems to be moderately expressed, but we did not see expression of *KXL1* and *KXL2*. We found a small band in *KXL1* (Figure 3), but since the band was much smaller than our expected band size of 454 base pairs, and also appeared in the negative control, we expect that this was due to primer dimerization. The expression of *KN* may imply that cell proliferation occurs in the petal, causing spurs to emerge. It at least indicates that cytokinesis and cell division is happening in some part of the petal.

STM1 and *STM2* are both genes that encode transcription factors maintaining cell indeterminacy in the

Primer	Sequence	Predicted Product Size
AqHistone4 Forward1+Reverse1	5'-AAG GCG TGG TGG TGT TAA GCG TAT CA - 3' 5'-GAA TTA CAA GAA AGT AGT AGA TCA GAA TCC AAC-3'	331 base pairs
AqSTM Forward + Reverse	5'-ATT ATC CAA GGC TCT TAG CTT G-3' 5'-CCG GTC AAA AGC ATC ACC AC-3'	308 base pairs
AqSTM2 Forward +Reverse	5'-TCT TCT CTG ATG ATG ATT CTG AAC AAA-3' 5'-CCT GAT GCG TAG GCC TCT TCC AAC T-3'	287 base pairs
AqKN Forward1 + Reverse1	5'-TCA ACA ACA ACA ACA GCA GCA G-3' 5'-TAA TTC CTC GCG ATA TTT TGC C-3'	473 base pairs
AqKXLI Forward + Reverse	5'-AAA GTG GAT CTG AGA TGA TGA GCG AT-3' 5'-GAT CGG AGA CTT GAA GAT TCT TGG CT-3'	454 base pairs
AqKXL2 Forward + Reverse	5'-CAA GGA ACT ACT GAA GGA AGT GGT G-3' 5'-GGC TGA AAC TCT GGA ACC TCT AAT TC-3'	434 base pairs
TCP4 Forward2 + Reverse2	5'-CCT GTT TAG GTT GGA CTC TAT GAG CT-3' 5'-GCA TCA GCC ATC TTT GTT GGT TTA CT-3'	196 base pairs
IPP2 Forward + Reverse	5'-CAG GTG AAG ACG GAC TGA AGT TTA A-3' 5'-CCA AGA CTG GAA AAA AGA CCA CAC-3'	182 base pairs
CYC Forward + Reverse	5'-GGG CAT TCT TAA AAT CAG CAA GGA TAA AGT GG-3' 5'-GAG GAC TGC TTA GAA CAT CCA CAA ACA CTC-3'	300 base pairs

Table 1. Primer Sequences and Size. RT-PCR used the listed forward and reverse primers to test the expression levels of each respective *KNOX* gene in *Aquilegia* petal spurs. Their full primer names, sequences, and predicted product size are shown above. *IPP2* is the positive control of the experiment.

apical meristem. Collani *et al.* never amplified *STM1* at any phase of petal development. Contrary to their results, we amplified a band of our expected size (308 base pairs) in our RT-PCR of *STM1* (Figure 3). We also amplified some larger bands. This was not what we expected, as we used the same primers to amplify *STM1* that Collani *et al.* used in their research. The multiple bands could be due to mis-splicing; *STM1* is one of several orthologs of the *Arabidopsis* locus in *Aquilegia*. The primers may not be as specific as we thought and might be amplifying more orthologs.

In addition, *CYC* was not expressed at all and this gene has been found to be responsible for promoting petal lobe growth and dorsoventral patterning in *Antirrhinum* (snapdragon). This implies that the dorsoventral differences seen in *Aquilegia* are promoted by a different transcription factor.

Discussion

Our results differed from Collani *et al.* in that while they found expression of *KN* and *STM1* in the inflorescence but not in petals, we saw expression of both of the transcription factors in our petal samples. One can hypothesize a reason for why this is the case. *STM* in general allows the meristem to produce a ready amount of stem cells. This low-intensity expression could be the cause of a longer cell proliferation phase. These results indicate that further work is necessary to elucidate whether *Aquilegia* petal spurs are dependent on *KNOX* genes.

To address these issues in mixed expression levels

from these *KNOX* genes, it may be beneficial to test *KNOX* expression in different phases of petal development. *CYC* may be activated in earlier or even later stages than we predicted. To investigate this possibility, RNA extraction of petals spurs in both the earlier and the later stages of development will be required.

We would also like to repeat Collani *et al.*'s *in situ* hybridization and examine gene expression localization of *STM1* in *Aquilegia* petals, also at differing stages of growth. *In situ* hybridization will help us investigate whether *HIS* and *TCP4* also have differing zones of expression within the petal.

Furthermore, gene expression studies do not confirm if the gene products are necessary for the morphology of the petal. Functional studies such as viral-induced knock-downs of *STM1* and *KN* would be useful to investigate whether impaired function of the protein products leads to petal spur defects.

It is clear that these genetic modules have extremely dynamic expression patterns. *KNOX* genes remain puzzling regulators of any plant, and to understand more about their function, more extensive tests must be performed that will allow us to obtain increasingly clear results.

Methods

Plant Material

All floral expression studies were performed using *Aquilegia coerulea* L. "Origami". Plants were grown in the greenhouses of the Dept. of Organismic and Evolutionary Biology, Harvard University.

RNA extraction and cDNA synthesis

RNA was extracted from petals using the Invitrogen Concert prep protocol for small scale extractions (Invitrogen, Carlsbad, CA, USA). cDNA was synthesized following the Invitrogen SuperScript III First-Strand Synthesis protocol (Invitrogen, Carlsbad, CA, USA).

RT-PCR

For each gene of interest, primers specific to that gene were mixed with distilled H₂O, Taq polymerase, cDNA, and 10x PCR buffer. The mix and cDNA was amplified for 25 cycles at 55 degrees Celsius. Amplification of the housekeeping control gene ISOPENTYL PYROPHOSPHATE: DIMETHYLALLYL PYROPHOSPHATE ISOMERASE2 (AqIPP2) was used as a positive control (Figure 4). See Table 1 for sequences and expected product size from all PCR primers.

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