Identification and Analysis of Jasmonate Pathway Genes in 
*Coffea canephora* (Robusta Coffee) by In Silico Approach

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**ABSTRACT**

Background: *Coffea canephora* is the commonly cultivated coffee species in the world along with *Coffea arabica*. Different pests and pathogens affect the production and quality of the coffee. Jasmonic acid (JA) is a plant hormone which plays an important role in plants growth, development, and defense mechanisms, particularly against insect pests. The key enzymes involved in the production of JA are lipoxygenase, allene oxide synthase, allene oxide cyclase, and 12-oxo-phytodienoic reductase. There is no report on the genes involved in JA pathway in coffee plants. **Objective:** We made an attempt to identify and analyze the genes coding for these enzymes in *C. canephora*. 

**Materials and Methods:** First, protein sequences of jasmonate pathway genes from model plant *Arabidopsis thaliana* were identified in the National Center for Biotechnology Information (NCBI) database. These protein sequences were used to search the web-based database Coffee Genome Hub to identify homologous protein sequences in *C. canephora* genome using Basic Local Alignment Search Tool (BLAST).

**Results:** Homologous protein sequences for key genes were identified in the *C. canephora* genome database. Protein sequences of the top matches were in turn used to search in NCBI database using BLAST tool to confirm the identity of the selected proteins and to identify closely related genes in species. The protein sequences from *C. canephora* database and the top matches in NCBI were aligned, and phylogenetic trees were constructed using MEGA6 software and identified the genetic distance of the respective genes. The study identified the four key genes of JA pathway in *C. canephora*, confirming the conserved nature of the pathway in coffee. The study expected to be useful to further explore the defense mechanisms of coffee plants. 

**Conclusion:** JA is a plant hormone that plays an important role in plant defense against insect pests. Genes coding for the 4 key enzymes involved in the production of JA viz., LOX, AOS, AOC, and OPR were identified in *C. canephora* (robusta coffee) by bioinformatic approaches confirming the conserved nature of the pathway in coffee. The findings are useful to understand the defense mechanisms of *C. canephora* and coffee breeding in the long run.

**Key words:** 12-oxo-phytodienoic reductase, allene oxide cyclase, allene oxide synthase, *Coffea canephora*, jasmonic acid pathway, lipoxigenase

**SUMMARY**

- JA is a plant hormone that plays an important role in plant defense against insect pests. Genes coding for the 4 key enzymes involved in the production of JA viz., LOX, AOS, AOC and OPR were identified and analyzed in *C. canephora* (robusta coffee) by in silico approach. The study has confirmed the conserved nature of JA pathway in coffee; the findings are useful to further explore the defense mechanisms of coffee plants.

**INTRODUCTION**

Jasmonic acid (JA) and its derivatives jasmonates (JAs) play a key role in plant metabolic processes and signaling systems when they are in stress and injured from pests.\(^1\)\(^-\)\(^4\) JAs, also called as oxylipins, are derived from fatty acid \(\alpha\)-linolenic acid through the formation of different intermediates, i.e., 13-hydroperoxy-9,11,15-octadecatrienoic acid by 13-lipoxygenase (LOX), 12-oxo-phytodienoic acid (OPDA) by allene oxide synthase (AOS) and allene oxide cyclase (AOC) in cytoplasm, and finally JA by OPDA reductase (OPR) through 3-oxo-2(2’\(Z\))-pentenyl) cyclopentane-1-octanoic acid by \(\beta\)-oxidation in peroxisomes [Figure 1].\(^1\)\(^-\)\(^4\) The synthesis and signaling process of JA and JAs as what we presently know is from studies on *Arabidopsis thaliana* and *Solanum lycopersicum*.\(^7\)\(^-\)\(^9\) Still, in many plant species including coffee, the jasmonate pathway is yet to be studied in detail. These plants have to be studied to know if they possess exactly the same biosynthetic pathway comprising similar enzymes that perform the same function in coffee as they do in model plants.

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the same biological role in their metabolism as in *Arabidopsis* and *Solanum*.

Coffee is one of the most important agricultural commodities; nearly 124 species are identified and exist around the world. However, only two species are considered commercially important, i.e., *Coffea arabica* L. and *Coffea canephora*, of which *C. arabica* is known for better cup quality compared to *C. canephora*. Between the two species, arabica is more sensitive to pathogens and pests (fungi, nematodes, and insects). In recent years, many coffee growers are shifting to robusta coffee cultivation due to high production costs, labor scarcity, and disease and pest management associated with arabica cultivation.

The objective of the study was to identify key genes of JA biosynthesis pathway in *C. canephora*, which is one of the parents of the tetraploid arabica coffee along with *Coffea eugenioides*. Recently, *C. canephora* genome is sequenced producing a high-quality draft genome of the species. The complete genome sequences of *C. canephora* are available in the web-based database of Coffee Genome Hub (http://coffee-genome.org/). The database served as a very useful resource in this study.

**MATERIALS AND METHODS**

**Databases**

GenBank Database of National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov) and database of Coffee Genome Hub (http://coffee-genome.org/) were used to identify the presence of proteins of the key enzymes involved in JA synthesis.

**Software**

Molecular evolutionary genetic analysis software MEGA6 (http://mega6.software.informer.com/) was used for sequence alignment and construction of phylogenetic trees.

**Identification of protein sequence of key genes of *Coffea canephora* involved in jasmonic acid biosynthesis**

Protein sequence of different genes involved in the JA synthesis was taken from the model plant *A. thaliana* using NCBI database. Those sequences were used for Basic Local Alignment Search Tool (BLAST) search and the Coffee Genome Hub database to identify the respective gene sequences in *C. canephora*. Among the matches in *C. canephora*, top match for each gene was taken and used for BLAST search against NCBI database. The top matches for *C. canephora* proteins involved in the JA pathway obtained in NCBI BLAST search were used for sequence alignment to obtain phylogeny trees using MEGA6 software.

**Construction of phylogenetic trees**

Around 14 top matches in NCBI database for each protein involved in the JA pathway and 1 top match from coffee genome database were taken from the BLAST results and transferred the sequence data to MEGA ALIGN tool in MEGA6 software. Using CLUSTALW alignment, the obtained sequences were aligned and by neighbor-joining method with bootstrap value 1000, the phylogenetic trees were constructed.

**RESULTS AND DISCUSSION**

The main enzymes that play a key role in the biosynthesis of JA are LOX, AOS, AOC, and OPR. The enzyme LOX is involved in the production of the intermediate compound 13-HPOT by adding oxygen to δ-linolenic acid at either C9 or C13 position, with 9S- or 13S-hydroperoxides. AOS and AOC are involved in the formation of 12-OPDA through unstable compound, an allene oxide. It is encoded by single gene in *Arabidopsis*, two genes in tomato. OPR belongs to a small family of related flavin-dependent oxidoreductases forms the JA through β-oxidation in peroxisome. The proteins of these enzymes were taken from model plant *A. thaliana* in NCBI database. Selected protein sequences of *A. thaliana* were used to identify homologous protein sequences in *C. canephora* genome from Coffee Genome Hub database using BLAST tool. We identified the matching protein sequences related to LOX, AOS, AOC, and OPR enzymes with similarities ranging from 2 to 17 [Table 1].

The same sequences were cross-checked again from coffee genome database with NCBI database and identified as unnamed protein products related to LOX, AOS, AOC, and OPR enzymes in

**Table 1: *Arabidopsis thaliana* protein sequences of key genes involved in jasmonic acid biosynthesis from National Center for Biotechnology Information used as query and *Coffea canephora* matches obtained in coffee genome hub database**

| Gene name | NCBI accession numbers of *Arabidopsis thaliana* protein sequence used | Number of matches to *Coffea canephora* proteins in coffee genome hub | Top match in coffee genome hub (locus and E value) |
|-----------|------------------------------------------------------------------------|---------------------------------------------------------------------|--------------------------------------------------|
| LOX       | AAA97315.1                                                             | 11                                                                  | Cc00 g30760, 0                                    |
| AOS       | NP_199079.1                                                            | 7                                                                   | Cc10 g03380, 0                                    |
| AOC       | CAC38763.1                                                             | 2                                                                   | Cc07 g08040, 2e-81                               |
| OPR       | CAB66143.1                                                             | 17                                                                  | Cc06 g12110, 0                                    |

OPR: OPDA reductase; OPDA: 12-oxo-phytodienoic acids; AOC: Allene oxide cyclase; AOS: Allene oxide synthase; LOX: Lipoxygenase; NCBI: National Center for Biotechnology Information
Table 2: Matches obtained for 12-oxophytodienoate reductase in *Coffea canephora* from coffee genome database. *Arabidopsis thaliana* protein sequence CAB66143.1 was used as query

| Gene name | $E$ | Query cover | Identity percentage | Locus ID |
|-----------|-----|-------------|---------------------|----------|
| 12-OPR 3–OPR3–complete | 0 | 97.44 | 75.32 | Cc06_g12110 |
| Putative 12-OPR 1–OPR11–complete | 8e-98 | 65.22 | 55.08 | Cc10_g09310 |
| 12-OPR 2–OPR2–complete | 5e-138 | 95.65 | 53.99 | Cc10_g09320 |
| 12-OPR 2–OPR2–complete | 2e-139 | 96.68 | 52.76 | Cc10_g09360 |
| Predicted protein–OPR11–modules | 7e-18 | 18.93 | 51.35 | Cc10_g09300 |
| Putative 12-OPR 1–OPR11–complete | 8e-131 | 95.65 | 51.32 | Cc10_g16500 |
| 12-OPR 1–OPR11–fragment | 6e-95 | 71.16 | 51.25 | Cc10_g09330 |
| 12-OPR 2–OPR2–complete | 8e-134 | 95.65 | 51.05 | Cc10_g09340 |
| Putative 12-OPR 11–OPR11–complete | 5e-110 | 86.96 | 50.88 | Cc06_g02480 |
| 12-OPR 1–OPR1–complete | 1e-129 | 96.68 | 50.79 | Cc10_g09350 |
| 12-OPR 2–OPR2–complete | 6e-128 | 95.65 | 48.35 | Cc10_g16490 |
| Putative 12-OPR 11–OPR11–complete | 6e-109 | 91.82 | 48.06 | Cc00_g31410 |
| Putative 12-OPR 11–OPR11–complete | 1e-106 | 122.76 | 47.82 | Cc10_g09290 |
| 12-OPR 2–OPR2–fragment | 1e-37 | 35.81 | 47.14 | Cc10_g16510 |
| Putative 12-OPR 11–OPR11–complete | 3e-100 | 134.78 | 46.97 | Cc10_g16520 |
| 12-OPR 1–OPR1–complete | 2e-103 | 95.65 | 44.92 | Cc09_g03820 |
| 12-OPR 1–OPR1–fragment | 5e-35 | 40.15 | 43.31 | Cc10_g16530 |

OPR: OPDA reductase; OPDA: 12-oxo-phytodienoic acid

Table 3: Matches obtained for allene oxide synthase in *Coffea canephora* from coffee genome hub database. *Arabidopsis thaliana* protein sequence NP_199079.1 was used as query

| Gene name | $E$ | Query cover | Identity percentage | Locus ID |
|-----------|-----|-------------|---------------------|----------|
| AOS, chloroplastic–CYP74A–complete | 0 | 99.42 | 66.41 | Cc10_g03580 |
| Hypothetical protein–CYP74A–missing_functional_completeness | 1e-44 | 22.59 | 64.10 | Cc00_g35230 |
| AOS, chloroplastic–CYP74A–complete | 0 | 89.56 | 63.81 | Cc10_g03570 |
| Hypothetical protein–CYP74A–missing_functional_completeness | 2e-42 | 22.59 | 62.39 | Cc05_g09930 |
| AOS–CYP74A2–complete | 1e-172 | 90.15 | 52.65 | Cc02_g18130 |
| 9-divinyl ether synthase–DES–complete | 7e-174 | 89.58 | 52.55 | Cc02_g18120 |
| Putative AOS, chloroplastic–CYP74A–complete | 2e-119 | 86.87 | 40.66 | Cc05_g03650 |

AOS: Allene oxide synthase

Table 4: Matches obtained for lipoxygenase in *Coffea canephora* genome from coffee genome hub database. *Arabidopsis thaliana* protein sequence AAF97315.1 was used as query

| Gene name | $E$ | Query cover | Identity percentage | Locus ID |
|-----------|-----|-------------|---------------------|----------|
| Linoleate 13S-lipoxygenase 3-1, chloroplastic–LOX3.1–complete | 0 | 87.61 | 73.41 | Cc00_g30760 |
| Linoleate 13S-lipoxygenase 3-1, chloroplastic–LOX3.1–fragment | 5e-69 | 19.96 | 63.24 | Cc05_g01710 |
| Lipoxigenase 6, chloroplastic–LOX6–complete | 0 | 93.42 | 54.62 | Cc11_g16680 |
| Linoleate 13S-lipoxygenase 3-1, chloroplastic–LOX3.1–complete | 0 | 99.56 | 49.58 | Cc02_g13400 |
| Linoleate 13S-lipoxygenase 2-1, chloroplastic–LOX2.1–complete | 0 | 81.69 | 47.81 | Cc01_g04060 |
| Linoleate 13S-lipoxygenase 1-1, chloroplastic–LOX1.1–fragment | 2e-20 | 12.94 | 45.11 | Cc00_g27370 |
| Probable linoleate 9S-lipoxygenase 5–LOX1.5–complete | 0 | 85.96 | 44.71 | Cc02_g33790 |
| Probable linoleate 9S-lipoxygenase 5–LOX1.5–complete | 0 | 85.96 | 44.71 | Cc02_g33800 |
| Probable linoleate 9S-lipoxygenase 5–LOX1.5–complete | 0 | 91.56 | 44.68 | Cc02_g33320 |
| Probable linoleate 9S-lipoxygenase 5–LOX1.5–complete | 0 | 92 | 44.25 | Cc02_g33780 |
| Linoleate 9S-lipoxygenase 5, chloroplastic–LOX5–complete | 0 | 94.74 | 43.62 | Cc03_g30580 |

LOX: Lipoxygenase

Table 5: Matches obtained for allene oxide cyclase in *Coffea canephora* from coffee genome hub database. *Arabidopsis thaliana* protein sequence CAC83763.1 was used as query

| Gene name | $E$ | Query cover | Identity percentage | Locus ID |
|-----------|-----|-------------|---------------------|----------|
| AOC 4, chloroplastic–AOC4–complete | 2e-81 | 74.81 | 64.77 | Cc07_g09040 |
| AOC 4, chloroplastic–AOC4–complete | 4e-89 | 100 | 55.73 | Cc11_g10540 |

AOC: Allene oxide cyclase

*C. canephora*. The maximum number of similarities obtained in coffee genome database was 17 for OPR [Table 2], 7 for AOS [Table 3] and 11 for LOX [Table 4]. The minimum number of similarities obtained was 2 for AOC [Table 5]. Top one similarity for each enzyme was taken as main source to confirm with NCBI by doing BLAST. Matches in NCBI database for each gene were taken for phylogenetic tree construction.

The evolutionary history was inferred using the neighbor-joining method. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) is shown next to the branches. The evolutionary distances were computed using the Poisson correction method and are in the units of the number of amino acid substitutions per site. The analysis involved...
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15 amino acid sequences. Evolutionary analyses were conducted in MEGA6, which was taken from coffee genome was showing bootstrap value 100. In addition to this, T. cacao and G. hirsutum also showed bootstrap value 100 [Figure 5]. The present study confirms the presence of genes involved in the synthesis of JA pathway in C. canephora, and further studies are needed to know the complete mechanism of these genes during signal transduction in resistance from pests compared to other coffee species.

CONCLUSION

JA pathway is reported to have mainly LOX, AOS, AOC, and OPR enzymes occurs in chloroplast and peroxisome, but there was no report of the presence of these enzymes in coffee plants. From the above study, we are concluding that the JA synthesis enzymes present in the pest-resistant coffee plant C. canephora and identified the close relatives of each gene in the NCBI database related to coffee plant. This study will be useful to understand the resistance mechanism in C. canephora, and this study is helpful to increase the resistance in other coffee plants such as C. arabica by artificial JA application in their cultivation.
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Conflicts of interest
There are no conflicts of interest.

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