COMPARATIVE PHYLOGENETIC ANALYSIS OF METHYLERYTHRITOL-4-PHOSPHATE (MEP) PATHWAY GENES IN ASTERACEAE FAMILY

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ABSTRACT: Stevioside glycosides (SGs) are a class of diterpene glycosides being produced by leaves of Stevia rebaudiana. Stevia is getting agricultural importance due to the production of natural sweetener steviol glycosides, also called as zero-calorie sweeteners. Earlier it was thought that steviol was synthesized from Kaurene via MVA pathway but later on it was concluded that precursors of steviol are actually synthesized via the plastid localized methlyerythritol 4-phosphate (MEP) pathway. In total, seventeen enzymes are used in this biosynthetic pathway. Each enzyme plays a significant role in the subsequent production of rebaudioside A and other SGs. In order to understand those enzymes functionally, understanding of those genes evolution basically in the Asteraceae family is highly warranted. In this study, the National Center for Biotechnology Information (NCBI) was searched to retrieve amino acid sequences that encode MEP pathway enzymes homologs. A number of MEP pathway enzymes’ amino acid sequences from the Asteraceae family were phyletically analyzed to light the way for the evolution characteristics of sweetener-producing plants homologs. The study is about to interpret the evolutionary relationships of Stevia rebaudiana with other plants in the same or different genus in the family Asteraceae.

INTRODUCTION: Isoprenoids are a class of natural products having important biological activities. All isoprenoids are constructed from two precursors, isopentenyl diphosphate (IPP) and its isomer dimethylallyl diphosphate (DMAPP). In the recent years, two most important discoveries in isoprenoid biosynthetic studies are the elucidation of a second pathway (the methlyerythritol phosphate (MEP) pathway) for isoprenoid biosynthetic and a modified mevalonate (MVA) pathway.

Isoprenoids are the largest and structurally diverse groups of natural products with more than 55,000 members that play key metabolic and regulatory roles in all life forms. Several monoterpenes, sesquiterpenes, and diterpenes produced as secondary metabolites are pheromones or defensive agents and are involved in many aspects of our life including flavor, fragrance, medicine and nutrition. Among that, a natural sweetening agent is Steviol Glycosides (SGs) which is being produced from Stevia rebaudiana. Stevia is getting agricultural importance due to the production of natural sweetener steviol glycosides, also called as zero-calorie sweeteners. The genus Stevia is comprised of approximately 200 species, which are distributed in North and South America. The phylogenetic relationships have been constructed using sequences of ITS and cpDNA.
and estimated the divergence times of the major clade of this genus. Results suggested that *Stevia* originated in Mexico 7.0–7.3 million years ago. Tissue-specific chemical analyses confirmed that SGs were accumulated in leaf cells but not in trichomes. Several steviol glycosides are produced by Stevia. Rebaudioside A is produced in the largest concentrations but possesses a bitter aftertaste. Whereas rebaudioside D is reported to have a superior flavor, but are produced in much smaller concentrations. The leaves of *Stevia rebaudiana* (Bertoni) is reported to accumulate at least eight steviol glycosides (SGs). Due to the differential glycosylation process, each SG possesses distinctive organoleptic properties. Stevioside is reported to be 143 times sweeter than sucrose on a weight basis, but Rebaudioside-A is 242 times sweeter. The aftertaste quality of Rebaudioside-A is better than stevioside because it is sweeter and less bitter. World Health Organization (WHO) has now recognized that stevioside is not genotoxic and has been assigned an acceptable daily intake of 0-2 mg SGs/kg body weight. Long back studies on Steviol concluded that it was synthesized from kaurene, via the mevalonate pathway (MVA) similar to the synthesis of several diterpenes.

**FIG. 1: MEP PATHWAY FOR PRODUCTION OF REBAUDIOSIDE A**

**TABLE 1: THE NAME OF THE GENES OF THESE PATHWAYS AND THEIR ABBREVIATION IS GIVEN BELOW WITH THE COMPARISON AT THE PROTEIN LEVEL TO THEIR ARABIDOPSIS ORTHOLOGS**

<table>
<thead>
<tr>
<th>S. no.</th>
<th>Enzymes</th>
<th>GenBank Acc no.</th>
<th>% Identity/Similarity to Acc no.**</th>
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<tbody>
<tr>
<td>1</td>
<td>DXS</td>
<td>AC143010.1</td>
<td>74/86 to Q38854</td>
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<tr>
<td>2</td>
<td>DXR</td>
<td>CAD22156.1</td>
<td>79/89 to AAF73140</td>
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<td>CMS</td>
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<td>4</td>
<td>CMK</td>
<td>ABB88838.3</td>
<td>70/80 to NP_180261.1</td>
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<td>5</td>
<td>MCS</td>
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<td>6</td>
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<td>8</td>
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<td>ABD92926.2</td>
<td>63/80 to F34802</td>
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<tr>
<td>10</td>
<td>CDPS*</td>
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<td>13</td>
<td>KAH*</td>
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<td>UGT85C2*</td>
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<td>UGT74G1*</td>
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<td>17</td>
<td>UGT76G1</td>
<td>ACT33422.1</td>
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(*for their absence in genbank databases those genes are excluded from our work.)

Deoxyxylulose-5-phosphate synthase (DXS), deoxyxylulose-5-phosphate reductoisomerase (DXR), 4-diphosphocytidyl-2-C-methyl-D-erythritol synthase (CMS), 4-diphosphocytidyl-2-C-methyl-D-erythritol kinase (CMK), 4-diphosphocytidyl-2-C-methyl-D-erythritol 2,4-cyclodiphosphate synthase (MCS), 1-hydroxy-2-methyl-2(E)-butenyl 4-diphosphate synthase (HDS) and 1-hydroxy-2-methyl-2(E)-butenyl 4-diphosphate reductase (HDR), geranylgeranyldiphosphate synthase (GGDSPS), copalyl diphosphate synthase (CDPS), kauren synthese (KS), kaurene oxidase (KO), kauroenoic acid 13-hydroxylase (KAH), UDP-glycosyltransferase (UGT).
Later it was demonstrated that the precursors of steviol are actually synthesized via the plastid localized methylerythritol 4-phosphate (MEP) pathway Fig. 1. 10 In total, seventeen enzymes are used in this biosynthetic pathway. Each enzyme plays a significant role in the subsequent production of rebaudioside A. In order to understand those enzymes functionally, understanding of those genes evolution basically in the Asteraceae family is highly warranted. The seventeen enzymes are enlisted in Fig. 1. 11

So in today’s scenario, effort is required in order to find out the superior varieties which may produce a range of SGs or knowledge about the biosynthetic (MEP) pathway genes to undergo for flux change through metabolic engineering or even finding the relationship of Stevia with other plantlets in the same or different genus in the family Asteraceae. The elucidation of biochemical pathways to study plant secondary metabolism has been a long term interest in the scientific community. The emergence of genomic resources like whole genome sequencing and tools like high throughput expressed sequence tags (ESTs) have accelerated the understanding of these metabolic pathways. 1

In this respect, to understand exact relationships between Stevia genus, other similar plants members in the family Asteraceae, Steviol glycoside content and composition, more predictable approaches are necessary. To gain insights into this, using the computational tools could enable the researchers to understand the physical and biochemical properties of proteins and genes involved. So the present study has been undertaken to find out the similarities between members of the Asteraceae family and to compare the MEP pathway genes in the Asteraceae family.

MATERIALS AND METHODS:
A) Phylogenetic Analysis:
Retrieval of the Protein Sequences of Enzymes & Screening of the Databases: Each gene of Stevia rebaudiana are searched by doing BLAST (http://www.ncbi.nlm.nih.gov/BLAST/) respect to the orthologs of Arabidopsis sp. Genbank accession number (as shown in Table 1 11 with the percentage of identity). The homology search of the genes was done by BLAST search tool of NCBI (https://www.ncbi.nlm.nih.gov/BLAST/) using blastp algorithm with an amino acid sequence of Asteraceae (taxid4210) family. Different amino acid sequences were downloaded from GenBank Accession number of different species of Asteraceae family 11.

Multiple Sequence Alignment: All protein sequences of genes were aligned using MUSCLE by MEGA software (version 7.0.21) to find out the similarity present among the sequences of the same family 12.

Phylogenetic Tree Construction: Phylogenetic analysis of the sequences was done by Molecular Evolutionary Genetics Analysis (MEGA) 13 software (version 7.0.21), using Neighbor-joining 14 method. Each node was tested using the bootstrap approach by taking 100. 15 The branch lengths were drawn to scale indicated.

B) Motifs’ Search:
Retrieval of the Protein Sequences of Enzymes & Screening of the Databases: Each gene of Stevia rebaudiana aare searched by doing BLAST (http://www.ncbi.nlm.nih.gov/BLAST/) respect to the orthologs of Arabidopsis sp. Genbank accession number (as shown in Table 1 with percentage of identity). The homology search of the genes was done by BLAST search tool of NCBI (https://www.ncbi.nlm.nih.gov/BLAST/) using blastp algorithm with amino acid sequence of Asteraceae (taxid4210) family. Different amino acid sequences were downloaded from genbank Accession number of different species of Asteraceae family 11.

Finding Motifs: Motif search is done by MEME-suite (http://meme-suite.org/) 17 of those protein sequences consequently by keeping one occurrence per sequence in select site distribution option and thirty-five in the option of select the numbers of motifs and other parameters are taken as customized in that page.

RESULTS AND DISCUSSION:
A) Phylogenetic Analysis: In this study, amino acid sequences of a total of eleven enzymes belonging to the MEP pathway were obtained from NCBI Genbank and analyzed. The neighbor-joining method was utilized to analyze the evolutionary history 14. MEGA7 was used to perform evolutionary analyses 13. According to the neighbor-joining algorithm, groups and subgroups
were defined per protein in phylogenetic tree. Convergence and divergence are two essential phylogenetic properties, which could be useful to identify the closely as well as distantly related groups of MEP pathway proteins. These findings suggest that all tested enzymes are conserved and all of the analyzed sequences evolved from a common ancestral enzyme. Each amino acid sequences of MEP pathway enzymes was first selected and screened, aligned and then subjected for phylogenetic analysis. So, a total number of eleven enzymes were taken and used for the final results processing. The results of each are showing and elaborating one after another.

No. (i) Deoxyxylulose-5-phosphate Synthase (DXS) Enzyme: The DXS enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to transketolase C super-family. From phylogenetic tree below based on the amino acid sequences of DXS enzyme, it is observed that Stevia rebaudiana and Helianthus annus are too closely related and diversified from the same origin. Similarly, crenenthum x morifolium and Taraxacum kok-saghyz are diversified from the same origin and are hence closely related to Cynara cardunculus var. scolymus related Atractylodes ianca are showing some similarity which means it is diversified from rest of species of Asteraceae family.

No. (ii) Deoxyxylulose-5-phosphate Reducto-Isomerase (DXR) Enzyme: The DXR enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to deoxyxylulose-5-phosphate reductoisomerase C super-family.

Similar sequences are randomly selected for multiple sequence alignment and phylogenetic analysis. Total eleven different species of Asteraceae family were selected from blastp result. For phylogenetic tree based on the protein sequences of DXR enzyme, Stevia rebaudiana is closely related to Taraxacum kok-saghyz and Helianthus annus; they are then more diversified to Atractylodes ianca and Argentinian adenophora. The both Chrysanthemum sp., Artemisia sp. and Achillea sp. diversified from each other from the same origin; which means they are distantly related from S. rebaudiana along with Tenacetum parthenium.

No. (iii) 4- diphosphocytidyl-2-C -methyl-D-Erythritol Synthase (CMS) Enzyme: The CMS enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to the Asteraceae family. The blast results depict that the enzyme belongs to glycosyl-transferase super-family.

Sequences above and are 99% were selected for multiple sequence alignment and phylogenetic analysis. Total five different species of Asteraceae family were selected from blastp result. For phylogenetic tree based on the protein sequence of CMS enzyme, Stevia rebaudiana is closely related to Taraxacum kok-saghyz then diversified with Cynara cardunculus var. scolymus. Both Helianthus annus and Artemisia annua are distantly related.

No. (iv) 4- diphosphocytidyl-2-C -methyl-D-Erythritol Kinase (CMK) Enzyme: The CMK enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to Ispe superfamily.

Multiple sequence alignment and phylogenetic analysis are done with selected five sequences. For phylogenetic tree based on the protein sequence of CMK enzyme, Stevia rebaudiana is closely related to Helianthus annus then to Taraxacum kok-saghyz then more diversified with Cynara cardunculus var. scolymus; Artemisia annua is distantly related.

No. (v) 4- diphosphocytidyl-2-C -methyl-D-Erythritol 2, 4-Cyclodiphosphate Synthase (MCS) enzyme: The MCS enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to MECDP synthase super-family.

Sequences which shows higher sequence similarity with six members of the family Asteraceae of MCS amino acid sequence of Stevia rebaudiana. Multiple sequence alignment and phylogenetic analysis are done with selected six sequences.
For phylogenetic tree based on the protein sequences of MCS enzyme, it is observed that Stevia rebaudiana is closely related to Ageratina adenophora; these two species are related to Taraxacum kok-saghyz and collectively these three species are related to Helianthus annus and then these four species are related to Cynara cardunculus var. scolymus; Artemisia annua is distantly related Stevia sp.

No. (vi) 1-hydroxy-2-methyl-2(E)-butenyl 4-Diphosphate Synthase (HDS) Enzyme: The HDS enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to GcpE protein super-family. Sequences above 80% were selected for multiple sequence alignment and phylogenetic analysis. Total five different species of Asteraceae family were selected from blastp result. For phylogenetic tree based on the protein sequences of HDS enzyme, it is observed that Stevia rebaudiana is closely related to Artemisia annua and these to are related with Helianthus annus. Taraxacum kok-saghyz and Cynara cardunculus var. scolymus are distantly related to Stevia sp.
No. (vii) 1-hydroxy-2-methyl-2(E)-butenyl 4-Diphosphate Reductase (HDR) Enzyme: The HDR enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to protein lytB and ispH super-family. Sequences above 99% were selected for multiple sequence alignment and phylogenetic analysis. Total six different species of asteraceae family were selected from blastp result. For phylogenetic tree based on the protein sequence of HDR enzyme, it is observed that Stevia rebaudiana is closely related to Tanacetum parthenium and Artemisia annua distantly related with Cynara cardunculus var. scolymus.

No. (viii) Geranyl-geranyl-diphosphate Synthase (GGDPS) Enzyme: The GGDPS enzyme of Stevia with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to protein isoprenoid biosynthesis class I super-family. Sequences above 100% identity were selected for multiple sequence alignment and phylogenetic analysis. Total of eight different species of Asteraceae family were selected from blastp result. For phylogenetic tree based on the amino acid sequence of GGDPS enzyme, it is observed that Stevia rebaudiana is distantly related to any other species of Asteraceae family.

No. (ix) Kaurene Synthase (KS) Enzyme: The KS enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to terpene synthase super-family.

No. (x) Kaurene Oxidase (KO) Enzyme: The KO enzyme of Stevia rebaudiana with a particular accession number was subjected for blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to terpene synthase super-family.

Sequences above 95% identity were selected for multiple sequence alignment and phylogenetic analysis. Total five different species of Asteraceae family were selected from blastp result. For phylogenetic tree based on the amino acid sequence of KO enzyme, it is observed here that Stevia is closely related with Helianthus annus and these both are diversified from Lactuca sativa; distantly related with Tanacetum cinerariifolium.
No. (xi) UDP-glycosyltransferase 76G1 (UGT 76G1) Enzyme: The UGT76G1 enzyme of *Stevia rebaudiana* with a particular accession number was subjected to blastp with respect to Asteraceae family. The blast results depict that the enzyme belongs to glycosyltransferase GTB type superfamily. Sequences above 98% identity were selected for multiple sequence alignment and phylogenetic analysis. Total five different species of Asteraceae family were selected from blastp result.

For phylogenetic tree based on the amino acid sequence of UGT76B1 enzyme, by different enzymes of species *Helianthus annus* selected from the alignment of the blastp result to allow the bootstrap process for tree construction. Here, we can see that UGT76B1 amino acid sequence is closely related to UGT76G1 of *Cynara cardunculus* var. scolymus but UGT76G1 of both *Stevia rebaudiana* and *Helianthus annus* are diversified from the same origin.

B) Motifs’ Search:

(i) For enzymes, *Deoxyxyulose-5-phosphate synthase* (DXS) & *Deoxyxyulose-5-phosphate reductoisomerase* (DXR) & *Geranyl-geranyl-diphosphate synthase* (GGDPS): Fig. 13A depicts the thirty-five motifs’ locations in each protein sequences of three enzymes with their consequent Genbank accession number with a p-value ranging from 1.58e-74 to 2.55e-119. Some of the motifs of these three enzyme sequences are shown below.

The five different motifs as raw are-

No.1 TGVGC MCIAC No.2 VHYDLWCRDY VHDDLPMADD No.3 FQVVDD DQVVDD
No.4 YYYGVP NWDGPK No.5 MMFKNEK EMFIDDK

In Fig. 13B some motifs which were found using MEME-suite; These five motifs are selected on the basis of E-value ranging from 1.0e+002 to 1.3e+004; varying with a length of six to twelve; the different colors indicate the presence of different amino acids in the protein sequences.

(ii) For enzymes *4-diphosphocytidyl-2-C-methyl-D-erythritol synthase* (CMS) & *4-diphosphocytidyl-2-C-methyl-D-erythritol kinase* (CMK): Fig. 14A depicts the thirty-five motifs’ locations on two protein sequences of CMS & CMK enzymes with a p-value ranging between 3.95e-134 to 5.25e-127 also along with their accession numbers.

The five different motifs as raw are-

No.1 QLVCVND EDVCIHD No.2 YTTSRMIRIQEI YCFGRGEEVKDI MAPLAA No.3 MSTMNA
No.4 QSSYTN QGNYRN No.5 DSFVPQ DSFANA

The motifs in Fig. 14B are found using MEME-suite; These five motifs are selected on the basis of E-value ranging from 1.0e+002 to 1.3e+004; varying with a length of six to twelve; the different colors indicate the presence of different amino acids in the protein sequences.

(iii) For Enzymes 1-hydroxy-2-methyl-(E)-butenyl 4-diphosphate reductase (HDR) & 1-hydroxy-2-methyl-(E)-butenyl 4-diphosphate synthase (HDS): The thirty-five motifs located on HDR & HDS enzymes’ protein sequences have a p-value ranging from 1.83e-159 to 3.33e-148.

The different motifs as raw are-

No.1 QRVNPNG IRIGPNG No.2 NNHFLS DIHFAP No.3 HPIRNQT AGIAIQT
No.4 VQGWN YGGWDS No.5 YYEYTW NCESTH
FIG. 13A: MOTIF LOCATIONS ON DXS, DXR & GGDPS ENZYMES’ PROTEIN SEQUENCES WITH THEIR ACCESSION NUMBER

FIG. 13B: 5 DIFFERENT MOTIFS OF DXS, DXR & GGDPS ENZYMES’ PROTEIN SEQUENCES

FIG. 14A: MOTIF LOCATIONS ON CMK, CMS ENZYMES’ PROTEIN SEQUENCES WITH THEIR ACCESSION NUMBER

FIG. 14B: 5 DIFFERENT MOTIFS OF CMK, CMS ENZYMES’ PROTEIN SEQUENCES
These five motifs are selected on the basis of E-value ranging from 1.1e+003 to 1.8e+001; varying with a length of six to seven Fig 15A and B.

(iv) For enzymes Kaurene oxidase (KO) & Kaurene synthase (KS): The thirty-five motifs located on KO & KS enzymes’ protein sequences have a p-value ranging from 1.04e-134 to 2.30e-144.

The different motifs as raw are-

<table>
<thead>
<tr>
<th>No.</th>
<th>Motif</th>
<th>p-value</th>
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<td>No.1</td>
<td>DTTWVAM</td>
<td>1.04e-134</td>
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<tr>
<td>No.2</td>
<td>DKAMVAM</td>
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</tr>
<tr>
<td>No.3</td>
<td>YSMKTG</td>
<td>2.30e-144</td>
</tr>
<tr>
<td>No.4</td>
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</tr>
<tr>
<td>No.5</td>
<td>QGYTVP</td>
<td></td>
</tr>
</tbody>
</table>

These five motifs are selected on the basis of E-value ranging from 1.1e+002 to 1.9e+001; varying with a length of six to eight Fig 16A and B.

(v) For enzymes UDP-glycosyltransferase 76G1 (UGT76G1) & 4-diphosphocytidyl-2-C-methyl-D-erythritol 2, 4-cyclodiphosphate synthase (MCS): The thirty-five motifs located on UGT76G1 & MCS enzymes’ protein sequences have a p-value ranging from 2.33e-117 to 5.41e-135.

The different motifs as raw are-

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<td>No.4</td>
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</tr>
<tr>
<td>No.5</td>
<td>ADQISCFT</td>
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</tr>
</tbody>
</table>

These five motifs are selected on the basis of E-value ranging from 1.0e+005 to 1.7e+003; varying with a length of six to eight Fig 17A and B.
CONCLUSION: In conclusion, amino acid sequence analysis of Methylerythritol-4-phosphate pathway proteins showed that these plants have been related together as they evolved together with conserved regions in sweetener production; there are some common species of Asteraceae family e.g., Helianthus annus is common in all the phylogenetic trees; the other common species are Cynara cardunculus var. scolymus and Taraxacum kok-saghyz. There are a bunch of motifs found in motifs’ search with two or three amino acids same and with a p-value ranging from 1.04e-134 to
5.41e-135. Motifs which are shown here mostly are with higher ranges from 1.0e+002 to 1.09e+001.

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REFERENCES:


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