Phyton (Austria)

16. XII. 1970

23 - 30

# Studies in the Identification of Compositae Taxa by Paper Chromatography

By

Krishna Mohan Madan DAKSHINI & PRITHIPAL SINGH \*)

#### With 1 Figure

Received September 25, 1969

# Introduction

Paper chromatographic techniques have recently been used for identification of various plant taxa (RILEY & BRYANT 1961; ELLISON et al. 1962; TORRES & LEVIN 1964; TURNER & MABRY 1964). However, for *Compositae*, one of the largest dicotyledonous angiosperm taxon, such data is not available. Further, *Compositae* are a well defined and a closely inter-related family in which "the taxa tend to be founded on slighter differences than in other families" (SMALL 1917). Prompted by these, the present chemotaxonomic investigation of *Compositae* of Delhi State (with special reference to the tribe *Heliantheae*) was undertaken to elicit information from biochemical compounds for identification purposes.

Organic compounds like, etheral oil, sesqueterpeulactone, diterpene, sterine, paraffine, saponine, carotinoids, wadelolactone, patulitrin, acetylenepolyene, alkaloids, cyanogen derivatives, aurones, chalcones, flavones, phenolic derivatives, sugars, organic acids, etc., have been reported to occur in *Compositae* taxa (PURI & SESHADRI 1954; BANNERJEE & SESHADRI 1956; GOVINDACHARI et al. 1956; SESHADRI & THAKUR 1960; HEGNAUER 1964). However, none of these compounds have been studied from the taxonomic viewpoint. BATE-SMITH 1962 demonstrated the taxonomic significance of phenolic constituents of plants, and indicated that *Compositae* like *Leguminosae* and unlike other *Campanulatae* have an exceptional number and diversity of phenolic compounds.

In the present investigation, therefore, the usefulness of the phenolics and their varied fluorescence types have been taken as the criterion for identification of 15 *Compositae* taxa.

<sup>\*)</sup> Dr. K. M. M. DAKSHINI, Department of Botany, University of Delhi, Delhi 7, India.

#### 24

We are grateful to Professor B. M. JOHRI for facilities and encouragement. Thanks are due to Dr. M. R. PARTHASARTHY \*) and Mr. R. K. TANDON for helpful suggestions.

# Materials and Methods

Aerial vegetative parts of 15 Compositae taxa (of tribe Heliantheae) occurring in and around Delhi, were collected during June to November 1968 and air-dried. The extraction of the phenolics from the dried plant tissue was done by acidic methanol. Whatman No. 1 chromatographic paper was used for separation and the chromatograms were developed by the descending method with BAW (*n*-Butanol:Glacial Acetic Acid:Water — 3:1:1). The developed chromatograms were scanned under UV with and without ammonia. Spots so developed (in UV, with and without NH<sub>3</sub>) were noted and numbered on their fluorescence types (Fig. 1), and their R<sub>t</sub>'s were calculated. The following abbreviations were used for denoting the colour and intensity of the spots: Br = brown; Bl = blue; Gr = green; V = violet; W = white; Yl = yellow; Fl fluorescent; P = pale; In = intense; Or = orange; R = red.

The arrangement of taxa in the figures and table follows that of "Flora of British India" (HOOKER 1882).

#### Results

Twenty-seven major spot types could be distinguished in 15 species studied. All taxa investigated showed spot No. 1 which fluoresced as Br under UV and did not change on fuming with  $NH_3$ . The  $R_f$  of this spot varied from 0,914 to 0,946 in different taxa (Table 1). The profile patterns of the taxa investigated are illustrated in Fig. 1. Based on the profile patterns of individual taxa, the species studied can be grouped in four categories:

Category A (6 to 12 spots in the profile; 2 or more spots showed changes in fluorescence with NH<sub>3</sub>): *Helianthus tuberosus*, *Glossocardia linearifolia*, *Bidens bipinnata*, *Bidens* sp., *Hymenatherum tenuifolium*.

Category B (only 3 spots in the profile; no change with  $NH_3$ ): Parthenium hysterophorus, Xanthium strumarium, Verbesina encelioides, Bidens biternata, Tridax procumbens.

Category C (4 to 6 spots in the profile; 1 or 2 spots show change in fluorescence with  $NH_3$ ): Sclerocarpus africanus, Blainvillea rhomboidea, Cosmos sulphureus, Gaillardia pulchella.

Category D (a characteristic aggregation of spot numbers 11, 12, 13 and 14): *Eclipta erecta*.

\*) Department of Chemistry, University of Delhi, Delhi 7, India.

### Category A

Helianthus tuberosus (9 spots), Glossocardia linearifolia (9 spots), Bidens bipinnata (9 spots), Bidens sp. (12 spots), and Hymenatherum tenuifolium (6 spots) showed similarity in the distributional pattern of the spots (Fig. 1). Each taxon, however, could be identified by changes observed after exposing the chromatograms to fumes of  $NH_3$ . In Helianthus tuberosus



Fig. 1. Uni-directional chromatographic patterns (with BAW) of aerial vegetative part extracts from 15 Compositae taxa. A = Parthenium hysterophorus, B = Xanthium strumarium, C = Eclipta erecta, D = Sclerocarpus africanus. E = Blainvillea rhomboidea, F = Helianthus tuberosus, G = Verbesina encelioides, H = Glossocardia linearifolia, I = Cosmos sulphureus, J = Bidens bipinnata, K = Bidens biternata, L = Bidens sp., M = Tridax procumbens, N = Hymenatherum tenuifolium, O = Gaillardia pulchella. (Bl = blue; Br = brown; Fl = fluorescent; I = intense; Or = orange; P = pale; R = red; V = violet).

and Glossocardia linearifolia, spot No. 4 was common and fluoresced as FlBl under UV and became InFlBl with  $NH_3$  fumes. In addition to spot No. 4 the profile of *H. tuberosus* had spot Nos. 19 ( $R_f 0.788$ ) and 21 ( $R_f 0.320$ ) which developed Yl fluorescence on exposure to  $NH_3$  fumes. Glossocardia linearifolia did not show spot Nos. 19 and 21 and was thus separated from *H. tuberosus*. Spot No. 4 did not occur in any other taxon of this category. The two species of Bidens (*B. bipinnata* and Bidens sp.) and Hymenatherum tenuifolium possessed spot Nos. 2 and 3. However, spot No. 2 fluorescing as

26

Table 1

 $R_f$  values and colours of spots developed in the chromatographic profile pattern  $UV: B_l = blue, Br = brown, Gr = green, V = violet, W = white, Yl = yellow, NH<sub>3</sub> are indicated in brackets.$ 

	Spot Types	Taxa						
Spot Nos.	Fluorescence under UV	Α	В	С	D	$\mathbf{E}$	$\mathbf{F}$	
1	Br	0.945	0.934	0.941	0.941	0.945	0.945	
2	FlGrBl					0.829		
3	FIBI					(InFlGrBl)	0.853	
4	FlBl					0.748 (InFlBl)	0.755 (InFlBl)	
5	PV				0.764			
6	FIPBIYI					0.622	0.611	
7	FlGrYl							
8	PV							
9	$\mathbf{PV}$						0.513	
10	FlW				0.480 (InFlW)			
11	VYl	0.382	0.377	0.408	L.	0.403	0.393	
12	InV	0.349	0.342	0.373	0.397			
13	InV			0.443 (InFlBl)				
14	FIBIV			0.375				
15	FIV			0.257				
16	Yl							
17	PBr							
18	BrYl						0.233	
19							0.788	
20	-						(11)	
21	-						0.320	
22	a						(¥1)	
23								
24	PBl							
<b>25</b>	InBr							
26	PYI							
27	PBrV							

27

Table	1
Lante	

of 15 Compositae taxa A-O (see explanation to Fig. 1). Fluorescence under Fl = fluorescent, P = pale, In = intense. – Colours of spots developed with

Taxa								
$\mathbf{G}$	$\mathbf{H}$	I	J	K	$\mathbf{L}$	$\mathbf{M}$	N	0
0.941	0.938	0.931	0.934	0.936	0.930	0.946	0.946	0.914
	0.829		0.787		0.810		0.80	
			(GrYl)		(GrYl)		(InFlGr	Bl)
			0.861		0.872		0.893	
			(Yl)		(GrBl)		(GrBl)	
	0.740							
	(InFlB	31)						
0.753		0.746				0.776		
		0.627						
		(InFlBI	Yl)					
	0.593							
0.567			0.554		0.578		0.563	0.543
	0.425		0.425		0.430			
	(InFlW	V)						
	0.373		0.363	0.376	0.383	0.404		
		0.253						
	0.283		0.261 (YIV)	0.266	0.30			
	0.229		()		0.263			
					0.774			
					(Yl)			
	0.655							
	(Yl)							
								0.070
								0.670
								(BIGr)
								0.383
			0 =00		0 =00			(GrYI)
			0.702		0.723			
					(GrBl)			
							0.127	

0.165

0.149

0.308 0.308

#### 28

FIGrBl became GrBl with  $NH_3$  in the two species of *Bidens*, while InFIGrBl in *Hymenatherum tenuifolium*. The two species of *Bidens* (*B. bipinnata* and *Bidens* sp.) were separated from each other by spot No. 3 which being common to both showed variations in fluorescence with  $NH_3$ ; in *B. bipinnata*, spot No. 3 changed from FIBl to Yl, whereas in *Bidens* sp., spot No. 3 changed from FIBl to GrBl. In addition, *Bidens* sp. showed spot No. 19 ( $R_f$  0.774) which developed Yl fluorescence with  $NH_3$ , and spot No. 24 ( $R_f$  0.723) which changed from PBl to GrBl with  $NH_3$ .

# Category B

Parthenium hysterophorus, Xanthium strumarium, Verbesina encelioides. Bidens biternata, and Tridax procumbens possessed 3 spots each. None of the spots showed any change on fuming with NH<sub>3</sub>. Individual taxa in this category could be delineated on their spot types, Rr values of the spots. and on the fluorescences of the starting point on the chromatogram. Thus, Parthenium hysterophorus and Xanthium strumarium which showed identical spot types (spot Nos. 1, 11, and 12) could be separated from each other on the R<sub>f</sub> values (0.945, 0.934; 0.382, 0.377; 0.349, 0.342 respectively) of the spots, and on the fluorescence of the starting point on the chromatogram. In Parthenium hysterophorus, the starting point fluoresced as PBr, while in Xanthium strumarium it was BrV. The above mentioned characteristic spots (Nos. 1, 11, and 12) of Parthenium and Xanthium were not found in the remaining taxa of this category. The other taxa could be identified by their spot types alone: Verbesina encelioides: Spot Nos. 1, 5, and 8; Bidens biternata: Spot Nos. 1, 12, and 16; Tridax procumbens: Spot Nos. 1, 5, and 12.

# Category C

Sclerocarpus africanus (4 spots), Blainvillea rhomboidea (6 spots), Cosmos sulphureus (4 spots), and Gaillardia pulchella (5 spots) showed similar distributional pattern of spots. Of these, Sclerocarpus africanus possessed spot No. 10 ( $R_f$  0.480) which fluoresced as W under UV and as InW with  $NH_3$ . This spot was absent in the other three taxa of this category. Spot No. 15 fluorescing as FIV was present in Cosmos sulphureus and Blainvillea rhomboidea but, whereas the former had 4 spots the latter had 6, and thus could be distinguished from one another. Gaillardia pulchella could be identified from other taxa of this category by spot Nos. 22 (R 0.670), 23 ( $R_f$  0.383), and 27 ( $R_f$  0.308). These spots were absent in other taxa.

# Category D

*Eclipta erecta* was identified by its profile which contained an aggregation of four spots, Nos. 11, 12, 13, and 14 (besides spot Nos. 1 and 15), which was not found in any other taxon investigated.

### Conclusions

Spot types and profile patterns in 15 species investigated showed marked variations. Also, each taxon had a characteristic profile pattern suggesting the reliability of the utility of biochemical compounds for taxonomic purposes. This supports the views of FAIRBROTHERS 1968, HAR-BOURNE 1967, and MIROV 1967 about the possibility of employing chemical constituents in helping to characterize, describe, and classify taxa. Further, the importance of present investigation lies in the fact that identification of the taxa could be made by studying the biochemical compounds present in the vegetative parts rather than the reproductive parts of the plants (PURI & SESHADRI 1954; BANNERJEE & SESHADRI 1956; GOVINDACHARI et al. 1956; SESHADRI & THAKUR 1960). It was possible to identify not only different genera, but also different species of the same genus. Thus by this method three species of *Bidens* investigated could be distinguished easily by their profile pattern. The data presented above suggest the need for further biochemical investigations of *Compositae* taxa.

## Summary

Fifteen Compositae taxa (belonging to tribe Heliantheae) were identified through paper chromatography of biochemical compounds extracted from vegetative parts. On the basis of profile pattern developed on the chromatograms, the taxa studied were grouped in four categories. Further identification of taxa within each category was possible through  $R_{\rm f}$  values and fluorescence of spots.

#### Literature Cited

- BANNERJEE N. R. & SESHADRI T. R. 1956. Isolation and constitution of patulitran. — Indian Acad. Sci. 44 A: 284—286.
- BATE-SMITH E. C. 1962. Phenolic constituents of plants and their taxonomic significance. I. Dicotyledons. J. linn. Soc., Bot. 58: 95-173.
- ELLISON W. L., ALSTON R. E. & TURNER B. L. 1962. Methods of presentation of crude biochemical data for systematic purposes with particular reference to the genus *Bahia* (*Compositae*). — Am. J. Bot. 49: 599—604.
- FAIRBROTHERS F. E. 1968. Chemosystematics with emphasis on systematic serology. In: Herwood V. H. (ed.) Modern Methods in Plant Taxonomy. — London.
- GOVINDACHARI T. R., NAGARAJAN K. & PAI B. R. 1956. Wadelolactone from *Eclipta erecta*. J. scient. ind. Res. 15 B: 664–665.
- HARBOURNE J. B. 1967. Comparative Biochemistry of the Flavonoids. London.
- HEGNAUER R. 1964. Chemotaxonomie der Pflanzen. Band 3. Basel und Stuttgart.

HOOKER J. D. 1882. Flora of British India. Vol. III. - London.

MIROV N. T. 1967. The Genus Pinus. - New York.

30

- PURI B. & SESHADRI T. R. 1954. Survey of anthocyanins. IV. Chromatographic study of yellow garden flowers, and constituents of coreopsin. — J. scient. ind. Res. 13 B: 321-325.
- RILEY H. P. & BRYANT T. R. 1961. The separation of nine species of the Iridaceae by paper chromatography. — Am. J. Bot. 48: 133-137.
- SESHADRI T. R. & THAKUR R. S. 1960. The colouring matter of the flowers of *Carthamus tinctorius*. - Curr. Sci. 29: 54.
- SMALL J. 1917. The origin and development of Compositae. I. History of classification of the Compositae. – New Phytol. 16: 157–177.
- TORRES A. M. & LEVIN D. A. 1964. A chromatographic study of Cespitose Zinnias. — Am. J. Bot. 51: 639-643.
- TURNER B. L. & MABRY T. J. 1964. Partition chromatography as applied to taxonomic problems in the *Asteraceae*. Taxon 13: 11-14.

# **ZOBODAT - www.zobodat.at**

Zoologisch-Botanische Datenbank/Zoological-Botanical Database

Digitale Literatur/Digital Literature

Zeitschrift/Journal: Phyton, Annales Rei Botanicae, Horn

Jahr/Year: 1970

Band/Volume: 14\_1\_2

Autor(en)/Author(s): Dakshini Krishna Mohdan Madan

Artikel/Article: <u>Studies in the Identification of Compositae Taxa by Paper</u> <u>Chromatography. 23-30</u>