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INTERNATIONAL UNION FOR THE PROTECTION OF NEW VARIETIES OF PLANTS

Geneva

DRAFT

SUNFLOWER

UPOV Code(s): HLNTS_ANN

Helianthus annuus L.

GUIDELINES

FOR THE CONDUCT OF TESTS

FOR DISTINCTNESS, UNIFORMITY AND STABILITY

prepared by experts from Hungary to be considered by the Technical Working Party for Agricultural Crops at its forty-ninth session, to be held in Saskatoon, Canada, from 2020-06-22 to 2020-06-26

Disclaimer: this document does not represent UPOV policies or guidance

Alternative names:*

Botanical name	English	French	German	Spanish
Helianthus annuus L.	Sunflower	Soleil, Tournesol	Sonnenblume	Girasol

The purpose of these guidelines ("Test Guidelines") is to elaborate the principles contained in the General Introduction (document TG/1/3), and its associated TGP documents, into detailed practical guidance for the harmonized examination of distinctness, uniformity and stability (DUS) and, in particular, to identify appropriate characteristics for the examination of DUS and production of harmonized variety descriptions.

ASSOCIATED DOCUMENTS

These Test Guidelines should be read in conjunction with the General Introduction and its associated TGP documents.

TG/81/7(proj.2) Sunflower, 2020-05-07 2

ТΑ	BLE O	F CONTENTS	PA
1.	SUBJE	CT OF THESE TEST GUIDELINES	<u>3</u>
2.	MATE	RIAL REQUIRED	.3
3.	METH	OD OF EXAMINATION	<u>3</u>
	3.1 3.2 3.3 3.4 3.5	Number of Growing Cycles Testing Place Conditions for Conducting the Examination Test Design Additional Tests	3 3 5
4.	ASSES	SSMENT OF DISTINCTNESS, UNIFORMITY AND STABILITY	_
	4.1 4.2 4.3	Distinctness Uniformity Stability	. 6
5.	GROU	PING OF VARIETIES AND ORGANIZATION OF THE GROWING TRIAL	. <u>8</u>
6.	INTRO	DUCTION TO THE TABLE OF CHARACTERISTICS	. <u>9</u>
	6.1 6.2 6.3 6.4 6.5	Categories of Characteristics States of Expression and Corresponding Notes Types of Expression Example Varieties Legend.	. <u>9</u> . <u>9</u> . <u>9</u>
7.		OF CHARACTERISTICS/TABLEAU DES CARACTÈRES/MERKMALSTABELLE/TABLA DE CTERES	. <u>12</u>
8.	EXPLA	NATIONS ON THE TABLE OF CHARACTERISTICS	. <u>24</u>
	8.1 8.2	Explanations covering several characteristics Explanations for individual characteristics	
9.	LITER	ATURE	. <u>33</u>
10.	TECH	NICAL QUESTIONNAIRE	. <u>34</u>

PAGE

1. <u>Subject of these Test Guidelines</u>

- 1.1 These Test Guidelines apply to all varieties of *Helianthus annuus* L.
- 1.2 In the case of ornamental varieties, in particular, it may be necessary to use additional characteristics or additional states of expression to those included in the Table of Characteristics in order to examine Distinctness, Uniformity and Stability.

2. <u>Material Required</u>

- 2.1 The competent authorities decide on the quantity and quality of the plant material required for testing the variety and when and where it is to be delivered. Applicants submitting material from a State other than that in which the testing takes place must ensure that all customs formalities and phytosanitary requirements are complied with.
- 2.2 The material is to be supplied in the form of seed.
- 2.3 The minimum quantity of plant material, to be supplied by the applicant, should be:

0.5 kg.

In the case of hybrid varieties, an additional 0.5 kg of each component (e.g. for a single hybrid, the female lines (male sterile line and maintainer line) and the male line) should be submitted. In the case of male sterile lines, an additional 0.5 kg of the maintainer line should be submitted.

The seed should meet the minimum requirements for germination, species and analytical purity, health and moisture content, specified by the competent authority. In cases where the seed is to be stored, the germination capacity should be as high as possible and should be stated by the applicant.

- 2.4 The plant material supplied should be visibly healthy, not lacking in vigor, nor affected by any important pest or disease.
- 2.5 The plant material should not have undergone any treatment which would affect the expression of the characteristics of the variety, unless the competent authorities allow or request such treatment. If it has been treated, full details of the treatment must be given.

3. <u>Method of Examination</u>

- 3.1 Number of Growing Cycles
- 3.1.1 The minimum duration of tests should normally be two independent growing cycles.
- 3.1.2 The two independent growing cycles should be in the form of two separate plantings.
- 3.1.3 The testing of a variety may be conducted when the competent authority can determine with certainty the outcome of the test.
- 3.2 Testing Place

Tests are normally conducted at one place. In the case of tests conducted at more than one place, guidance is provided in TGP/9 "Examining Distinctness".

- 3.3 Conditions for Conducting the Examination
- 3.3.1 The tests should be carried out under conditions ensuring satisfactory growth for the expression of the relevant characteristics of the variety and for the conduct of the examination.
- 3.3.2 The optimum stage of development for the assessment of each characteristic is indicated by a number in the Table of Characteristics. The stages of development denoted by each number are described in Chapter 8.
- 3.4 Test Design
- 3.4.1 Each test should be designed to result in a total of at least 40 plants, which should be divided between at least 2 replicates.
- 3.4.2 The design of the tests should be such that plants or parts of plants may be removed for measurement or counting without prejudice to the observations which must be made up to the end of the growing cycle.
- 3.5 Additional Tests

Additional tests, for examining relevant characteristics, may be established.

- 4. Assessment of Distinctness, Uniformity and Stability
- 4.1 Distinctness
- 4.1.1 General Recommendations

It is of particular importance for users of these Test Guidelines to consult the General Introduction prior to making decisions regarding distinctness. However, the following points are provided for elaboration or emphasis in these Test Guidelines.

To assess distinctness of hybrids, the parent lines and the formula may be used according to the following recommendations:

(i) description of parent lines according to the Test Guidelines;

(ii) check of the originality of the parent lines in comparison with the variety collection, based on the characteristics in Chapter 7, in order to identify similar parent lines;

(iii) check of the originality of the hybrid formula in relation to the hybrids in the variety collection, taking into account the most similar lines; and

(iv) assessment of the distinctness at the hybrid level for varieties with a similar formula.

Further guidance is provided in documents TGP/9 "Examining Distinctness" and TGP/8 "Trial Design and Techniques Used in the Examination of Distinctness, Uniformity and Stability".

4.1.2 Consistent Differences

The differences observed between varieties may be so clear that more than one growing cycle is not necessary. In addition, in some circumstances, the influence of the environment is not such that more than a single growing cycle is required to provide assurance that the differences observed between varieties are sufficiently consistent. One means of ensuring that a difference in a characteristic, observed in a growing trial, is sufficiently consistent is to examine the characteristic in at least two independent growing cycles.

4.1.3 Clear Differences

Determining whether a difference between two varieties is clear depends on many factors, and should consider, in particular, the type of expression of the characteristic being examined, i.e. whether it is expressed in a qualitative, quantitative, or pseudo-qualitative manner. Therefore, it is important that users of these Test Guidelines are familiar with the recommendations contained in the General Introduction prior to making decisions regarding distinctness.

4.1.4 Number of Plants or Parts of Plants to be Examined

Unless otherwise indicated, for the purposes of distinctness, all observations on single plants should be made on 36 plants or parts of plants taken from each of 36 plants and any other observations made on all plants in the test, disregarding any off-type plants.

4.1.5 Method of Observation

The recommended method of observing the characteristic for the purposes of distinctness is indicated by the following key in the Table of Characteristics (see document TGP/9 "Examining Distinctness", Section 4 "Observation of characteristics"):

MG: single measurement of a group of plants or parts of plants MS: measurement of a number of individual plants or parts of plants VG: visual assessment by a single observation of a group of plants or parts of plants VS: visual assessment by observation of individual plants or parts of plants

Type of observation: visual (V) or measurement (M)

"Visual" observation (V) is an observation made on the basis of the expert's judgment. For the purposes of this document, "visual" observation refers to the sensory observations of the experts and, therefore, also includes smell, taste and touch. Visual observation includes observations where the expert uses reference points (e.g. diagrams, example varieties, side-by-side comparison) or non-linear charts (e.g. color charts). Measurement (M) is an objective observation against a calibrated, linear scale e.g. using a ruler, weighing scales, colorimeter, dates, counts, etc.

Type of record: for a group of plants (G) or for single, individual plants (S)

For the purposes of distinctness, observations may be recorded as a single record for a group of plants or parts of plants (G), or may be recorded as records for a number of single, individual plants or parts of plants (S). In most cases, "G" provides a single record per variety and it is not possible or necessary to apply statistical methods in a plant-by-plant analysis for the assessment of distinctness.

In cases where more than one method of observing the characteristic is indicated in the Table of Characteristics (e.g. VG/MG), guidance on selecting an appropriate method is provided in document TGP/9, Section 4.2.

4.2 Uniformity

- 4.2.1 It is of particular importance for users of these Test Guidelines to consult the General Introduction prior to making decisions regarding uniformity. However, the following points are provided for elaboration or emphasis in these Test Guidelines:
- 4.2.2 These Test Guidelines have been developed for the examination of [type or types of propagation] varieties. For varieties with other types of propagation the recommendations in the General Introduction and document TGP/13 "Guidance for new types and species", Section 4.5 "Testing Uniformity" should be followed.
- 4.2.3 The assessment of uniformity for open-pollinated should be according to the recommendations for cross-pollinated varieties in the General Introduction.
- 4.2.4 The assessment of uniformity for hybrid varieties depends on the type of hybrid and should be according to the recommendations for hybrid varieties in the General Introduction.
- 4.2.5 Where the assessment of a hybrid variety involves the parent lines, the uniformity of the hybrid variety should, in addition to an examination of the hybrid variety itself, also be assessed by examination of the uniformity of its parent lines.
- 4.2.5 For the assessment of uniformity of inbred lines, a population standard of 2% and an acceptance probability of at least 95% should be applied. In the case of a sample size of 36 plants, 2 off-types are allowed. In addition, the same population standard and acceptance probability should apply for the assessment of uniformity regarding out-crosses and isogenic male fertile plants in a male sterile line. For the assessment of uniformity of single hybrids, a population standard of 5% with an acceptance probability of at least 95% should be applied. In the case of a sample size of 36 plants, 4 off-types are allowed. For three-way hybrids and open-pollinated varieties, the variability within the variety should not exceed the variability of comparable varieties already known.

- 4.2.6 If enzyme electrophoresis is used for testing distinctness, the same population standard and the same acceptance probability as for other characteristics should be applied. All plants within an inbred line with one locus or more loci being heterozygous with one allele in each locus coming from the inbred line (e.g. AX) should be considered out-crosses. All other cases of heterozygosity as well as cases where one foreign allele is present in one locus with homozygous status should be considered off-types.
- 4.3 Stability
- 4.3.1 In practice, it is not usual to perform tests of stability that produce results as certain as those of the testing of distinctness and uniformity. However, experience has demonstrated that, for many types of variety, when a variety has been shown to be uniform, it can also be considered to be stable.
- 4.3.2 Where appropriate, or in cases of doubt, stability may be further examined by testing a new seed stock to ensure that it exhibits the same characteristics as those shown by the initial material supplied.
- 4.3.3 Where appropriate, or in cases of doubt, the stability of a hybrid variety may, in addition to an examination of the hybrid variety itself, also be assessed by examination of the uniformity and stability of its parent lines.

5. <u>Grouping of Varieties and Organization of the Growing Trial</u>

- 5.1 The selection of varieties of common knowledge to be grown in the trial with the candidate varieties and the way in which these varieties are divided into groups to facilitate the assessment of distinctness are aided by the use of grouping characteristics.
- 5.2 Grouping characteristics are those in which the documented states of expression, even where produced at different locations, can be used, either individually or in combination with other such characteristics: (a) to select varieties of common knowledge that can be excluded from the growing trial used for examination of distinctness; and (b) to organize the growing trial so that similar varieties are grouped together.
- 5.3 The following have been agreed as useful grouping characteristics:
 - (a) Leaf: intensity of green color (characteristic 2)
 - (b) Leaf: blistering (characteristic 3)
 - (c) Time of beginning of flowering (characteristic 12)
 - (d) Ray floret: color (characteristic 18)
 - (e) Disk flower: pro-duction of pollen (characteristic 21)
 - (f) Only inbred lines: Plant: natural height (characteristic 26)
 - (g) Only hybrids and open-pollinated varieties: Plant: natural height (characteristic 27)
 - (h) Plant: branching (characteristic 28)
 - (i) Seed: color (characteristic 38)
 - (j) Seed: stripes on margin (characteristic 39)
 - (k) Seed: stripes be-tween margins (characteristic 40)
- 5.4 Guidance for the use of grouping characteristics, in the process of examining distinctness, is provided through the General Introduction and document TGP/9 "Examining Distinctness".

6. <u>Introduction to the Table of Characteristics</u>

- 6.1 Categories of Characteristics
- 6.1.1 Standard Test Guidelines Characteristics

Standard Test Guidelines characteristics are those which are approved by UPOV for examination of DUS and from which members of the Union can select those suitable for their particular circumstances.

6.1.2 Asterisked Characteristics

Asterisked characteristics (denoted by *) are those included in the Test Guidelines which are important for the international harmonization of variety descriptions and should always be examined for DUS and included in the variety description by all members of the Union, except when the state of expression of a preceding characteristic or regional environmental conditions render this inappropriate.

6.2 States of Expression and Corresponding Notes

- 6.2.1 States of expression are given for each characteristic to define the characteristic and to harmonize descriptions. Each state of expression is allocated a corresponding numerical note for ease of recording of data and for the production and exchange of the description.
- 6.2.2 In the case of qualitative and pseudo-qualitative characteristics (see Chapter 6.3), all relevant states of expression are presented in the characteristic. However, in the case of quantitative characteristics with 5 or more states, an abbreviated scale may be used to minimize the size of the Table of Characteristics. For example, in the case of a quantitative characteristic with 9 states, the presentation of states of expression in the Test Guidelines may be abbreviated as follows:

State	Note
small	3
medium	5
large	7

However, it should be noted that all of the following 9 states of expression exist to describe varieties and should be used as appropriate:

State	Note
very small	1
very small to small	2
small	3
small to medium	4
medium	5
medium to large	6
large	7
large to very large	8
very large	9

6.2.3 Further explanation of the presentation of states of expression and notes is provided in document TGP/7 "Development of Test Guidelines".

6.3 Types of Expression

An explanation of the types of expression of characteristics (qualitative, quantitative and pseudoqualitative) is provided in the General Introduction.

6.4 Example Varieties

Where appropriate, example varieties are provided to clarify the states of expression of each characteristic.

TG/81/7(proj.2) Sunflower, 2020-05-07 8

6.5 Legend

		English		françai	S	deutsch español		Example Varieties Exemples Beispielssorten Variedades ejemplo	Note/ Nota
1	2	3	4	5	6	7	•	•	
	Name of characteristics in English		Nom o caract frança	tère en	Name des Merkmals auf Deutsch	Nombre del carácter en español			
	states of expressio			types	d'expression	Ausprägungsstufen	tipos de expresión		

1 Characteristic number

2	(*)	Asterisked characteristic	– see Chapter 6.1.2
3	Type of expression QL QN PQ	Qualitative characteristic Quantitative characteristic Pseudo-qualitative characteristic	– see Chapter 6.3 – see Chapter 6.3 = – see Chapter 6.3
4	Method of observation (and type MG, MS, VG, VS	of plot, if applicable)	– see Chapter 4.1.5
5	(+)	See Explanations on the Table of	of Characteristics in Chapter 8.2
6	(a)-(b)	See Explanations on the Table of	of Characteristics in Chapter 8.1

7 Growth stage key See Explanations on the Table of Characteristics in Chapter 8

7. <u>Table of Characteristics/Tableau des caractères/Merkmalstabelle/Tabla de caracteres</u>

		English			français	deutsch	español	Example Varieties Exemples Beispielssorten Variedades ejemplo	Note/ Nota
1.		QN	VG			10	•		
		Seedling: anthocyanin coloration of hypocotyl							
		absen	t or very weak					T0954LM	1
		weak						OB724	2
		mediu	m					99D40R	3
		strong						F7AW1MOA	4
		very s	trong					Kisvárdai	5
2.	(*)	QN	VG		(a), (b)	51-55			
		Leaf: i green	intensity of color						
		light						F5DN3MA, T0243HG	3
		medium						H11050R	5
		dark						13013	7
3.	(*)	QN	VG		(a), (b)	51-55		-	
		Leaf:	blistering						
		ahsen	t or very weak					F5DN3MA	1
		weak						F7AX2JA, IR79DMR	3
		mediu	 m					HA89, IB1088DMR	5
		strong						TRC2342	7
		very s							9
4.	(*)		VG	(+)	(a), (b)	51-55	-		
		i	serration						
			ed or very fine					99D40R	1
		fine 						IR79DMR	3
		mediu						HA89, TRC2342	5
		coarse						PB1458DMR	7
		very c	oarse						9

Example Varieties English français deutsch español Note/ Exemples Nota Beispielssorten Variedades ejemplo 5. VG QN (+) (a), (b) 53-55 Leaf: shape in cross section RT9513 1 concave flat PH5002R 2 3 convex 6. PQ VG 53-55 (a), (b) (+) Leaf: shape of distal part lanceolate FR810RM1 1 2 lanceolate to narrow triangular narrow triangular RT0976 3 4 medium triangular RT9513 broad triangular BT0835 5 6 short acuminate 7 broad triangular to SF9074MA rounded medium acuminate 8 rounded 9 7. (*) QN VG (+) (a), (b) 53-55 Leaf: auricules 37025 1 none or very small T0954LM small 3 medium IR79DMR 5 F6AH6MO, HA89 7 large 9 very large 8. VG 53-55 QN (+) (a), (b) Leaf: wings none or very weakly T0954LM 1 expressed 2 weakly expressed F7AW1MOA strongly expressed 13013 3

	English		français	deutsch	español	Example Varieties Exemples Beispielssorten Variedades ejemplo	Note/ Nota
9. (*)	QN VG	(+)	(a), (b)	53-55	•	•	
-	Leaf: angle of lowest lateral veins						
	acute					T0860LM	1
	right angle or nearly right angle					F7AW1MOA	2
	obtuse					TFC3767B	3
10.	QN VG	(+)	(a), (b)	53-55	•	•	
	Leaf: height of tip of blade compared to insertion of petiole						
	very low						1
	low						2
	medium					99D40R	3
	high					T0954LM	4
	very high						5
11. (*)	QN MS/VG		(a), (b)	55-57	•	-	
	Leaf: size						
	small					PH5002R	3
	medium					LC1093, OB724	5
	large					F5DN3MA	7
12. (*)	QN MG/MS	(+)	(a)	61			
	Time of beginning of flowering						
	very early					PHA283	1
	early					T0860LM	3
	medium					H11050R, RHA274	5
	late					RT7710	7
	very late					Kisvárdai, LGR27	9
13.	QN VG	(+)	(a)	63-65		•	
	Ray floret: attitude in relation to head						
	right angle					T0954LM	1
	right angle to horizontal						2
	horizontal					T0833HG	3

français **Example Varieties** Note/ English deutsch español Exemples Nota Beispielssorten Variedades ejemplo 14. VG PQ (a) 63-65 Ray floret: dis-position flat HA89, IR79DMR 1 longitudinal recurved PH5002R 2 F5DN3MA 3 undulated strongly recurved to back of head 4 15. QN VG (a) 63-65 Flower: density of ray florets T0954LM 1 very sparse medium 99D40R, HA89 3 OB724 5 very dense 16. QN MS/VG (+) (a) 63-65 Ray floret: width very narrow T0954LM 1 2 HA850, OB724 narrow 3 broad very broad 4 17. MS/VG 63-65 QN (a) Ray floret: length short BT0835 3 SF9074MA 5 medium 7 T0954LM long 18. (*) PQ VG (+) (a) 63-65 Ray floret: color yellowish white RHA381 1 F7AW1MOA 2 light yellow medium yellow RT7710 3 4 orange yellow U0881BG orange OB724, P211R 5 6 purple reddish brown 7

Example Varieties English français deutsch español Note/ Exemples Nota Beispielssorten Variedades ejemplo 19. PQ VG (a) 63-65 Disk flower: color yellow STR226, TRC2342 1 2 orange F7AW1MOA, HA89 3 purple 20. VG QN (+) (a) 63-65 Disk flower: an-thocyanin colora-tion of stigma SF9074MA absent or very weak 1 2 RT7710 weak medium R6ST2MI, TRC2342 3 F7AW1MOA 4 strong Kisvárdai 5 very strong 21. (*) QL VG (a) 63-65 Disk flower: pro-duction of pollen F7AW1MOA, HA89 1 absent present IR79DMR, RHA274 9 22. VG 63-65 QN (+) (a) Bract: shape clearly elongated T0954LM 1 neither clearly elongated nor clearly rounded IR79DMR 2 3 clearly rounded IB1088DMR MS/VG 23. QN (a) 63-65 (+) Bract: length of tip short IB1088DMR 3 medium HA89, T0954LM 5 long U0881BG 7 24. VG (a) 63-65 QN Bract: intensity of green color of outer side light T0243HG 1 medium T0954LM 3 RT8711 5 dark

		English		français	deutsch	español	Example Varieties Exemples Beispielssorten Variedades ejemplo	Note/ Nota
25.	QN	VG		(a)	69-73			
-		: attitude in on to head						
	not en slightl	nbracing or very y embracing					HA89, RT0976	1
	slightl	y embracing					F7AW1MOA	2
	strong	ly embracing					RT9513	3
26. (*)	QN	MS		(a)	69-73			
	<u>Only i</u> Plant:	inbred lines: : natural height						
	very s	hort					FR810RM1	1
	short						OB724	3
	mediu	ım					U0881BG	5
	tall						R6ST2MI	7
	very ta	all					31G03	9
27. (*)	QN	MS			69-73			
	open- variet	<u>hybrids and</u> pollinated ies: Plant: al height						
	very s	hort					Antonil	1
	short						GK Milia	3
	mediu	Im					Sumiko	5
	tall						Marley	7
_	very ta	all					Kisvárdai	9
28. (*)	QL	VG		(a)	69-89			
	Plant	branching						
	absen	ıt					HA89, OB724	1
	prese	nt					RHA274, T0954LM	9
29. (*)	PQ	VG	(+)	(a)	69-89			
	Plant: type of branching							
	only b	asal						1
	predo	minantly basal						2
	overal	11					H11050R	3
	predo	minantly apical					RHA274, T0954LM	4
	only a	pical					TRC2342	5

Note/ **Example Varieties** English français deutsch español Exemples Nota Beispielssorten Variedades ejemplo 30. VG QN 69-89 Plant: position of highest lateral head to central head R6ST2MI 1 below 2 same level T0954LM 3 above 99D40R 31. (*) QN VG 80-89 (a) Stem: attitude straight U0881BG 1 2 slightly curved strongly curved F7EW2MIA 3 4 over turned 32. (*) QN VG (a) 80-89 (+) Head: attitude horizontal RT8711 1 2 inclined RT0976 3 vertical half-turned down U0881BG 4 F5DN3MA 5 turned down over turned 6 33. (*) QN MS/VG 80-89 (a) Head: size small RT0976 3 medium BT0835, HA89 5 7 large F5DN3MA 34. (*) PQ VG 85-87 (+) (a) Head: shape of grain side strongly concave 1 2 weakly concave R5PG6MJ RT8711 3 flat HA89, R6ST2MI 4 weakly convex strongly convex F5DN3MA 5 TRC3398R 6 deformed

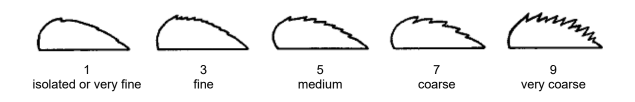
		English		français	deutsch	español	Example Varieties Exemples Beispielssorten Variedades ejemplo	Note Nota
35.	QN	MS/VG		(a)	99			
	Seed:	size						
	very s	mall					PHA283	1
	small						TRC2342	3
	mediu	Im					HA89, OB724	5
	large						FT2603, Kisvárdai	7
	very la	arge						9
36. (*)	PQ	VG	(+)	(a)	99	-		
	Seed:	shape						
	elonga	ated					BT0835	1
		w ovoid					H11050R	2
	broad						F7AW1MOA, HA89	3
	rounded						,	4
37.	QN	MS/VG		(a)	99	-		1
•	Seed: thickness relative to width							
	very tł	hin					RHA801	1
	thin							3
	mediu	ım					F7AW1MOA	5
	thick						85C11R, F7AX2MA	7
	very tł	hick						9
38. (*)	PQ	VG	(+)	(a)	99	_		
	Seed:	color						
	white						Labud	1
	purple							2
	light b						IR79DMR	3
	medium brown						H11050R	4
	dark b	prown					B0644LM	5
	light g	Irey						6
		ım grey					RT9513	7
	dark g	jrey						8
	black						HA89, T0954LM	9

		English		français	deutsch	español	Example Varieties Exemples Beispielssorten Variedades ejemplo	Note Nota
39. (*)	QN	VG	(+)	(a)	99			
	Seed: marg	: stripes on in		•				
	none expre	or very weakly ssed					T0954LM	1
	weakl	y expressed					OB724	2
	strong	gly expressed					HA89, U0881BG	3
40. (*)	QN	VG	(+)	(a)	99		•	
	Seed: marg	: stripes be-tween ins						
	none expre	or very weakly ssed					T0954LM	1
	weakl	y expressed					LGR27	2
	strong	gly expressed					HA89, U0881BG	3
41. (*)	PQ	VG		(a)	99		•	
•	Seed	color of stripes	Ī	•				
	white						U0881BG	1
	browr	1					F1164LM	2
	grey						99D40R	3
	black							4

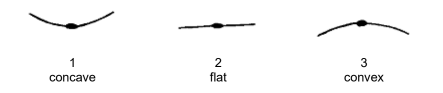
- 8. <u>Explanations on the Table of Characteristics</u>
- 8.1 Explanations covering several characteristics

Characteristics containing the following key in the Table of Characteristics should be examined as indicated below:

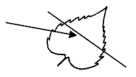
- (a) All observations should be made on the main stem.
- (b) All observations on the leaf should be made on fully developed leaves at the 2/3 height of the plant.
- 8.2 Explanations for individual characteristics
- Ad. 4: Leaf: serration



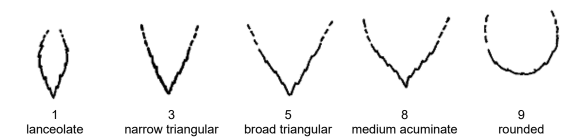
Ad. 5: Leaf: shape in cross section



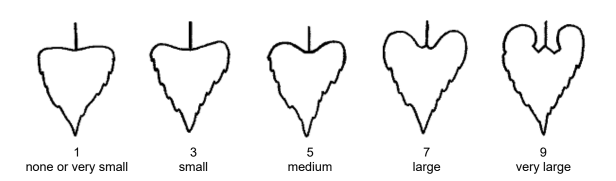
Cross section:



Ad. 6: Leaf: shape of distal part

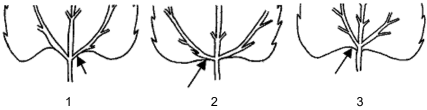


Ad. 7: Leaf: auricules



Ad. 8: Leaf: wings

(parenchym at base of lateral veins)



none or very weakly expressed

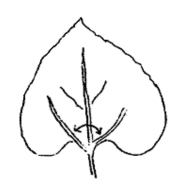
weakly expressed

strongly expressed

Ad. 9: Leaf: angle of lowest lateral veins



acute

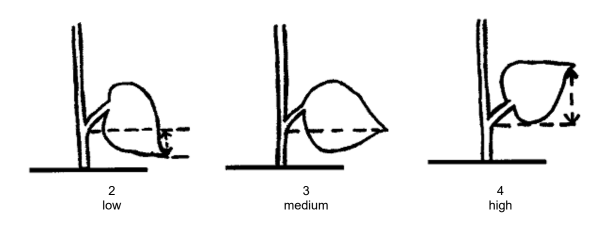


2 right angle or nearly right angle



3 obtuse

Ad. 10: Leaf: height of tip of blade compared to insertion of petiole



Ad. 12: Time of beginning of flowering

Time of flowering is reached when 50% of the plants have at least one extended ray floret.

Ad. 13: Ray floret: attitude in relation to head







Ad. 16: Ray floret: width





very narrow

2 narrow



3 broad



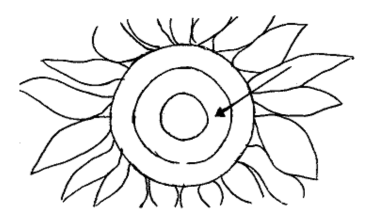
very broad

Ad. 18: Ray floret: color

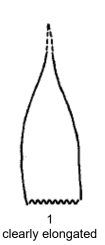
If more than one color, only the color covering the biggest surface is considered.

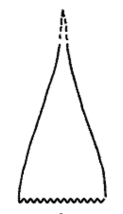
Ad. 20: Disk flower: an-thocyanin colora-tion of stigma

The anthocyanin coloration should be recorded on the stigma just after the pollen appears at the top of the anthers.

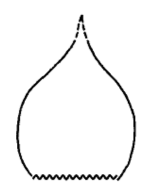


Ad. 22: Bract: shape



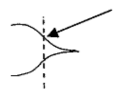


2 neither clearly elongated nor clearly rounded



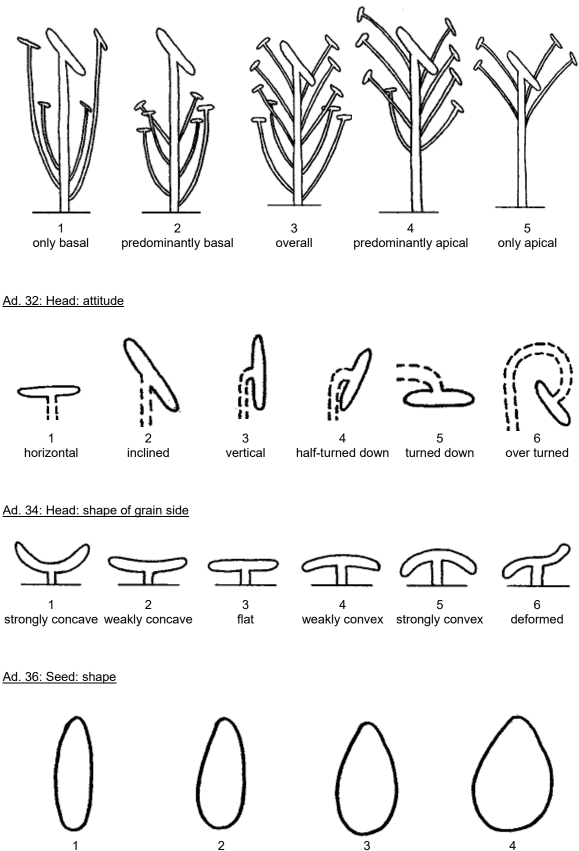
3 clearly rounded

Ad. 23: Bract: length of tip



Tip begins where the direction of curving changes

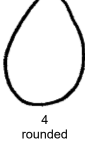
Ad. 29: Plant: type of branching



elongated

narrow ovoid

broad ovoid



Ad. 38: Seed: color

The main color of the seed is the color with the largest area. In case of doubt which is the largest area, the darkest color is the main color.

Ad. 39: Seed: stripes on margin



Ad. 40: Seed: stripes be-tween margins



8.3 Growth stage of Helianthus annuus L. adopted to the BBCH (Meier U., 1997) scale applicable to individual plant

Code Description

Principal growth stage 0: Germination

- 00 Dry seed (achene)
- 01 Beginning of seed imbibition
- 03 Seed imbibition complete
- 05 Radicle emerged from seed
- 06 Radicle elongated, root hairs developing
- 07 Hypocotyl with cotyledons emerged from seed
- 08 Hypocotyl with cotyledons growing towards soil surface
- 09 Emergence: cotyledons emerge through soil surface

Principal growth stage 1: Leaf development1

- 10 Cotyledons completely unfolded
- 12 2 leaves (first pair) unfolded
- 14 4 leaves (second pair) unfolded
- 15 5 leaves unfolded
- 16 6 leaves unfolded
- 17 7 leaves unfolded
- 18 8 leaves unfolded
- 19 9 or more leaves unfolded
- (Stem elongation may occur earlier than stage 19; in this case continue with the principal stage 3)

Principal growth stage 3: Stem elongation

- 30 Beginning of stem elongation
- 31 1 visibly extended internode
- 32 2 visibly extended internodes
- 33 3 visibly extended internodes
- 3. Stages continuous till . . .
- 39 9 or more visibly extended internodes

Principal growth stage 5: Inflorescence emergence

- 51 Inflorescence just visible between youngest leaves
- 53 Inflorescence separating from youngest leaves, bracts distinguishable from foliage leaves
- 55 Inflorescence separated from youngest foliage leaf
- 57 Inflorescence clearly separated from foliage leaves
- 59 Ray florets visible between the bracts; inflorescence still closed

Principal growth stage 6: Flowering

- 61 Beginning of flowering: ray florets extended, disc florets visible in outer third of inflorescence
- Disc florets in outer third of inflorescence in bloom (stamens and stigmata visible)

65 Full flowering: disc florets in middle third of inflorescence in bloom (stames and stigmata visible)

67 Flowering declining: disc florets in inner third of inflorescence in bloom (stames and stigmata visible)

69 End of flowering: most disc florets have finished flowering, ray florets dry or fallen **Principal growth stage 7: Development of fruit**

- 71 Seeds on outer edge of the inflorescence are grey and have reached final size
- 73 Seeds on outer third of the inflorescence are grey and have reached final size
- 75 Seeds on middle third of the inflorescence are grey and have reached final size
- 79 Seeds on inner third of the inflorescence are grey and have reached final size

Principal growth stage 8: Ripening

80 Beginning of ripening: seeds on outer third of anthocarp black and hard. Back of anthocarp still green

- 81 Seeds on outer third of anthocarp dark and hard. Back of anthocarp still green
- 83 Dark of anthocarp yellowish-green, bracts still green. Seeds about 50% dry matter

Seeds on middle third of anthocarp dark and hard. Back of anthocarp yellow, bracts brown edged. Seeds about 60% dry matter

87 Physiological ripeness: back of the anthocarp yellow. Bracts marbled brown. Seeds about 75–80% dry matter

89 Fully ripe: seeds on inner third of anthocarp dark and hard. Back of anthocarp brown. Bracts brown. Seeds about 85% dry matter

Principal growth stage 9:

- 92 Over ripe, seeds over 90% dry matter
- 97 Plant dead and dry
- 99 Harvested product

9. <u>Literature</u>

ASFIS, GEVES, GNIS: "Description des géniteurs et variétés de tournesol", édition 2000 (English, French, Spanish) ASFIS, 44, rue du Louvre, 75001 Paris, FR

Meier U., 1997: Growth stages of mono- and dicotyledonous plants: BBCH-Monograph. Wien Federal Biological Research Center for Agriculture and Forestry, Blackwell Wissenschafts-Verlag, Berlin, DE.

Miller J.F.: "Update on Inheritance of Sunflower Characteristics," USDA - ARS, Northern Crop Science Laboratory, Fargo, North Dakoto 58105, USA

10. <u>Technical Questionnaire</u>

TECHN		QUESTIONNAIRE	Page {x} of {y}	Reference Number:
				Application date: (not to be filled in by the applicant)
			ECHNICAL QUESTIONN	AIRE n for plant breeders' rights
1.	Subjec	t of the Technical Question	naire	
	1.1	Botanical name	Helianthus annuus L.	
	1.2	Common name	Sunflower	
2.	Applica	ant		
	Name	Γ		
	Addres	s		
	Teleph	one No.		
	Fax No	o. [
	E-mail	address		
	Breede applica	er (if different from [nt)		
3.	Propos	ed denomination and breed	ler's reference	
	Propos (if avai	ed denomination		
	Breede	er's reference		

TECHNICAL C	QUESTIONNAIRE	Page {x} of {y}		Reference Numbe	er:
#4. Informa	4. Information on the breeding scheme and propagation of the variety				
4.1	Breeding scheme				
Variety	resulting from:				
4.1.1	Crossing				
(a)	controlled cross				[]
	(please state parent variety)				
	() x		()
	female parent			male parent	
(b)	partially known cross				[]
	(please state known parent	variety(ies))			
	() x		()
	female parent			male parent	
(c)	unknown cross				[]
4.1.2	Mutation (please state parent variety) -				[]
4.1.3	Discovery and development (please state where and who		dev	veloped)	[]
4.1.4	Other (Please provide details)				[]

TECHNICAL (QUESTIONNAIRE	Page {x} of {y}	Reference Number:	
4.2 4.2.1	Method of propagating Other (Please provide details)	-	[]	

	Characteristics of the variety to be indicated (th characteristic in Test Guidelines; please mark t		ling
	Characteristics	Example Varieties	Note
5.1 (2)	Leaf: intensity of green color		
.,	light	F5DN3MA, T0243HG	3 [
	medium	H11050R	5 [
	dark	13013	7 [
5.2 (3)	Leaf: blistering		
.,	absent or very weak	F5DN3MA	1 [
	weak	F7AX2JA, IR79DMR	3 [
	medium	HA89, IB1088DMR	5 [
	strong	TRC2342	7 [
	very strong		9 [
5.3 (12)	Time of beginning of flowering		
. ,	very early	PHA283	1 [
	early	T0860LM	3 [
	medium	H11050R, RHA274	5 [
	late	RT7710	7 [
	very late	Kisvárdai, LGR27	9 [
5.4 (18)	Ray floret: color		
	yellowish white	RHA381	1 [
	light yellow	F7AW1MOA	2 [
	medium yellow	RT7710	3 [
	orange yellow	U0881BG	4 [
	orange	OB724, P211R	5 [
	purple		6 [
	reddish brown		7 [
5.5 (21)	Disk flower: pro-duction of pollen		
	absent	F7AW1MOA, HA89	1 [
	present	IR79DMR, RHA274	9 [

	Characteristics	Example Varieties	Note
5.6 (26)	Only inbred lines: Plant: natural height		
. ,	very short	FR810RM1	1[]
	short	OB724	3[]
	medium	U0881BG	5[]
	tall	R6ST2MI	7[]
	very tall	31G03	9[]
5.7 (27)	Only hybrids and open-pollinated varieties: Plant: na height	atural	
	very short	Antonil	1[]
	short	GK Milia	3[]
	medium	Sumiko	5[]
	tall	Marley	7[]
	very tall	Kisvárdai	9[]
5.8 (28)	Plant: branching		
	absent	HA89, OB724	1[]
	present	RHA274, T0954LM	9[]
5.9 (38)	Seed: color		
	white	Labud	1[]
	purple		2[]
	light brown	IR79DMR	3[]
	medium brown	H11050R	4[]
	dark brown	B0644LM	5[]
	light grey		6[]
	medium grey	RT9513	7[]
	dark grey		8[]
	black	HA89, T0954LM	9[]
5.10 (39)	Seed: stripes on margin		
	none or very weakly expressed	T0954LM	1[]
	weakly expressed	OB724	2[]
	strongly expressed	HA89, U0881BG	3[]
5.11 (40)	Seed: stripes be-tween margins		
	none or very weakly expressed	T0954LM	1[]
	weakly expressed	LGR27	2[]
	strongly expressed	HA89, U0881BG	3[]

TECHNICAL QUESTIONNAIRE	Page {x} of {y}	Reference Number:					
6. Similar varieties and differences from	these varieties						
from the variety (or varieties) which, to the	Please use the following table and box for comments to provide information on how your candidate variety differs from the variety (or varieties) which, to the best of your knowledge, is (or are) most similar. This information may help the examination authority to conduct its examination of distinctness in a more efficient way.						
Denomination(s) of Characteristic variety(ies) similar to your your candidate		e the expression of Describe the expression acteristic(s) for the the characteristic(s)					
Example							
Comments:							

TECHI	NICAL Q	UESTIONNAIRE	Page {x} of {y}	Reference Number:	
#7. 7.1					
	help to Yes	distinguish the variety?	No	[]	
7.2		please provide details) ere any special conditions for	growing the variety or con	ducting the examination?	
	Yes (If yes,	[] please provide details)	No	[]	
7.3	Other i	nformation			

TECI		AL QUESTIONNAIRE	Page (v) of (v)	Poforor	ice Number:	
		L QUESTIONNAIRE	Page {x} of {y}	Kelelel		
8.	Autho	prization for release				
	(a)	Does the variety require environment, human and		elease under legisl	ation concerning t	the protection of the
		Yes []	No []		
	(b)	Has such authorization b	een obtained?			
		Yes []	No []		
	If the	answer to (b) is yes, pleas	e attach a copy of the a	uthorization.		
9. In	formati	on on plant material to be e	examined or submitted	or examination		
9.2 chara	s and stocks, The pl acterist underg	e expression of a characte disease, chemical treatme scions taken from different ant material should not tics of the variety, unless to one such treatment, full de your knowledge, if the plan	ent (e.g. growth retard growth phases of a tre have undergone any he competent authoritie etails of the treatment n	ants or pesticides e, etc. treatment which v s allow or request nust be given. In th), effects of tissu would affect the such treatment. his respect, please	expression of the lf the plant material
	(a)	Microorganisms (e.g	. virus, bacteria, phytop	lasma)	Yes []	No []
	(b)	Chemical treatment	(e.g. growth retardant, _l	oesticide)	Yes []	No []
	(c)	Tissue culture			Yes []	No []
	(d)	Other factors			Yes []	No []
	Ple	ase provide details for whe	ere you have indicated '	yes".		
10.	l he	ereby declare that, to the b	est of my knowledge, th	e information prov	ided in this form is	s correct:
	Ар	plicant's name				
			L			
	Się	gnature		Date	e	

Common Sunflower, 2020-05-07

34

ANNEX

Part I Introduction

The following Annex contains a list of characteristics derived by using electrophoresis and a description of the method to be used. UPOV decided to place these characteristics in an Annex to the Test Guidelines, thereby creating a special category of characteristic, because the majority of the UPOV member States is of the view that it is not possible to establish distinctness solely on the basis of a difference found in a characteristic derived by using electrophoresis. Such characteristics should therefore only be used as a complement to other differences in morphological or physiological characteristics. UPOV reconfirms that these characteristics are considered useful but that they might not be sufficient on their own to establish distinctness. They should not be used as a routine characteristic but at the request or with the agreement of the applicant of the candidate variety.

	Charac	Part II Characteristics Derived by Using Electrophoresis				
Nr.	Characteristic	States of expression	Example varieties	Note		
42	Allele expression at locus Me1	Genotype 2/2	IB1088DMR	1		
		Genotype 4/4	SF9074MA	2		
		Genotype 2/4	Sumiko	3		
43	Allele expression at locus Pgd1	Genotype 2/2	IB1088DMR	1		
		Genotype 4/4	SF9074MA	2		
		Genotype 2/4	Sumiko	3		
44	Allele expression at locus Pgi2	Genotype 2/2	IB1088DMR	1		
		Genotype 4/4	SF9074MA	2		
		Genotype 2/4	GK Petrus CLP	3		
45	Allele expression at locus Shdh1	Genotype 2/2	IB1088DMR	1		
		Genotype 4/4		2		
		Genotype 2/4	Marley	3		
46	Allele expression at locus Pgm4	Genotype 2/2		1		
		Genotype 4/4	IB1088DMR	2		
		Genotype 2/4	GK Petrus CLP	3		

Common Sunflower, 2020-05-07

35

Part III

Description of the Method to be Used

Description of the SGE Method for the Analysis of Isoenzymes from Helianthus annuus L.

1. Number of seedlings per test :

 For checking formula: 10 seedlings each of inbred lines

4 seedlings of single hybrids 10 seedlings of three-way hybrids

• For distinctness, uniformity and stability test:

at least 40 seedlings for inbred lines, hybrids and open-pollinated varieties

2. Apparatus and equipment

Any suitable horizontal electrophoresis system can be used, provided that the gels can be kept at 4° C. A gel thickness of 10 mm is recommended. The power supply used should be capable of delivering constant voltage output.

3. Chemicals

All chemicals should be of 'Analytical Reagent' grade or better.

- 3.1 <u>Chemicals for enzyme extraction:</u> Tris- (hydroxymethyl) aminomethane (Tris) Hydrochloric acid β-Mercaptoethanol
- 3.2 <u>Chemicals for electrophoresis</u> Bromophenol blue Citric acid monohydrate L-Histidine Starch hydrolysed, for electrophoresis, (Sigma S-4501 or equivalent)
- 3.3 Chemicals for staining enzymes

95% Ethanol Ethylenediamine tetra-acetic acid, disodium salt (EDTA Na2) D-Fructose 6-phosphate, disodium salt α-D-Glucose 1-phosphate, monohydrate, disodium salt Glucose 6-phosphate dehydrogenase (Sigma G5885) Hydrochloric acid (HCl) Magnesium chloride hexahydrate (MgCl2, 6H2O) DL-Malic acid, monosodium salt Dimethylthiazol diphenyl tetrazolium (MTT) β-Nicotinamide adenine dinucleotide phosphate (NADP) Nitro-blue tetrazolium (NBT) 6-phosphogluconic acid, trisodium salt dihydrate Phenazine methosulfate (PMS) Shikimic acid Sodium hvdroxide (NaOH) Tris- (hydroxymethyl) aminomethane (Tris)

Common Sunflower, 2020-05-07

36

4. Solutions

- 4.1 Extraction solution: 0.1M Tris HCI (pH 7.2) + 0.2 % 2-mercaptoethanol (v/v).
- 4.2. Electrophoresis buffers
- 4.2.1 Buffers for SGE pH 6.5
- 4.2.1.1 Stock solution: 0.364 M L-histidine-citrate

50.44 g L-histidine 8.34 g Citric acid monohydrate made up to 1 l with de-ionised water

4.2.1.2 Running buffer: 0.072 M L-histidine-citrate pH 6.5 (Stock solution diluted 1 in 5)

400 ml stock solution (4.2.1.1) made up to 2 l with de-ionised water

4.2.1.3 Gel buffer: 0.024 M L-histidine-citrate (Stock solution diluted 1 in 15)

80 ml stock solution (4.2.1.1) made up to 1200 ml with de-ionised water

- 4.2.2 Buffers for SGE pH 5.7
- 4.2.2.1 Running buffer: 0.067 M L-histidine-citrate pH 5.7:

20.18 g L-histidine 8.34 g Citric acid monohydrate made up to 2 l with de-ionised water

4.2.2.2 Gel buffer: 0.011 M L-histidine-citrate (Running buffer diluted 1 in 6):

100 ml running buffer (4.2.2.1) made up to 1200 ml with de-ionised water

4.2.2.3 Bromophenol blue solution:

50 mg bromophenol blue dissolved in 100 ml de-ionised water

- 4.3 Staining solutions
- 4.3.1 Stock solutions
- 4.3.1.1 1 M Tris-HCl pH 7.5

121.1 g Tris, made up to 1 I with de-ionised water and adjusted to pH 7.5 with 50 % HCI

4.3.1.2 1 M Tris-HCl pH 8.5

121.1 g Tris, made up to 1 I with de-ionised water and adjusted to pH 8.5 with 50 % HCI

4.3.1.3 MTT solution

1.0 g MTT made up to 100 ml with de-ionised water

4.3.1.4 NBT solution

1.0 g NBT made up to 100 ml with de-ionised water

Common Sunflower, 2020-05-07

37

4.3.1.5 PMS solution

200 mg PMS made up to 100 ml with de-ionised water

4.3.1.6 MgCl2 solution

10 g Magnesium chloride hexahydrate made up to 100 ml with de-ionised water

4.3.1.7 Sodium malate solution

2.5 g DL-malic acid made up to 50 ml with de-ionised water and adjusted to pH 8.0 with 1M NaOH.

4.3.2 Staining solutions

4.3.2.1 ME staining solution

100 ml 0.1 M Tris HCl, pH 7.5 (4.3.1.1 diluted 1 in 10) 4 ml Sodium malate solution (4.3.1.7.) 1 ml NBT solution (4.3.1.4.) 1 ml PMS solution (4.3.1.5.) 1,8 ml MgCl2 solution (4.3.1.6.) 17.5 mg NADP

4.3.2.2 PGD + PGI staining solution

100 ml 0.1 M Tris HCl, pH 7.5 (4.3.1.1. diluted 1 in 10) 100 mg D-Fructose 6-phosphate Na2 salt 60 mg 6-Phosphogluconic acid Na3 salt 10 mg NADP 1 ml MTT solution (4.3.1.3.) 1.5 ml PMS solution (4.3.1.5.) 1 ml MgCl2 solution (4.3.1.6.) 40 units of Glucose-6-phosphate dehydrogenase (SIGMA G 5885) To stain PGI only, do not include 6-phosphogluconic acid. To stain PGD only, do not include either fructose 6-phosphate disodium salt or glucose 6-phosphate dehydrogenase.

4.3.2.3 ShDH staining solution

100 ml 0.2 M Tris HCl, pH 8.5 (4.3.1.2 diluted 1 in 5) 50 mg shikimic acid 1 ml MTT solution (4.3.1.3) 1.25 ml PMS solution (4.3.1.5) 12 mg NADP

4.3.2.4 PGM staining solution

100 ml 0.1 M Tris HCl, pH 8.5 (4.3.1.2. diluted 1 in 10) 150 mg α -D-Glucose 1-phosphate 1H2O, Na2 salt 150 mg EDTA, Na2 10 mg NADP 1.5 ml MTT solution (4.3.1.3) 1.ml PMS solution (4.3.1.5) 4 ml MgCl2 solution (4.3.1.6) 40 units of Glucose 6-phosphate dehydrogenase

Common Sunflower, 2020-05-07

38

5. Procedure

5.1. Enzyme extraction

Seedlings are grown on moistened germination paper, at 25°C, in darkness, for 2 to 3 days. Seed coats are removed and cotyledons are crushed at 4°C, with a pestle in 1.5 ml microtubes containing 300 μ l extraction buffer (4.1). The extracts can be stored at -30°C or at -80°C.

5.2 Preparation of the gel

Prepare the gels the day before migration. To make two 12.5 % starch gels (18 x 18 x 1 cm) the following is required: 128 g starch are mixed in 1020 ml gel buffer (4.2.1.3 or 4.2.2.2) in a 1000 ml Büchner flask and heated at 78°C. The mixture is degassed with a water jet aspirator for 30 seconds. The gels are poured into gel moulds as described in the user's manual of the equipment used. The formation of air bubbles should be avoided. The gels are allowed to cool at room temperature for 45 min, then placed in a refrigerator for 1 h. The gels are wrapped with polyethylene film for overnight storage. and cooled to 4°C for 1 h before migration.

5.3 Electrophoresis

5.3.1 Each electrode tank is filled with the appropriate volume of running buffer (4.2.1.2 or 4.2.2.1) pre-cooled to 4° C. The polyethylene film is lifted up and two transversal slits are cut in the gel 3 cm and 4 cm from the edge (cathode side) of the mould.

The 1 cm gel slice is removed and the extracts are loaded as follows:

The enzyme extracts are thawed from 5.1, and absorbed on a filter paper wick (1.5 mm x 20 mm, Whatman N $^{\circ}$ 3). The wicks are inserted into the gel, tightly against the first slit. One wick soaked with bromophenol blue solution (4.2.2.3) (migration dye marker) is placed on each side of the gel.

The gel slice is cautiously replaced. Each gel is covered with polyethylene film.

The two gels, with the extracts on the cathodal side, are placed on the two electrode buffer tanks, in a refrigerated cabinet at 4°C.

The electrophoresis is carried out at 4°C, towards the anode. After 15 min of migration at the first voltage, the wicks are removed and the voltage is increased. Constant voltage should be maintained during each phase.

The electrophoretic conditions are indicated in the following table.

Buffer systems	Constant voltage	Distance run by bromophenol blue	Duration of migration
Histidine citrate pH 5.7	260 V for 15 min then 290 V	13 cm	5 h
Histidine citrate pH 6.5	240 V for 15 min then 280 V	11 cm	5 h

SGE at pH 5.7 should be used for detecting ME, PGD and PGI. The isoenzymes PGM and ShDH should be analysed by SGE pH 6.5.

5.4 Enzyme staining

After switching off the current, the gel is cut horizontally in 1 mm thick slices with a very fine steel wire or a fishing line. The upper slice is discarded. Individual gel slices are stained by incubation at 37°C, in darkness in the following solutions:

for ME:	solution 4.3.2.1,	incubation time: 15 h
for PGD and PGI:	solution 4.3.2.2,	incubation time: 1 h
for SHDH:	solution 4.3.2.3,	incubation time: 1 h
for PGM:	solution 4.3.2.4,	incubation time: 1/2 h

Common Sunflower, 2020-05-07

39

After staining the gel slices are rinsed in de-ionised water and fixed in 40% ethanol solution. The following procedures for long time storing can be successfully used: e.g. drying of the gels between two cellophane sheets soaked in a 5% glycerol solution, or storing in sealed polyethylene bags.

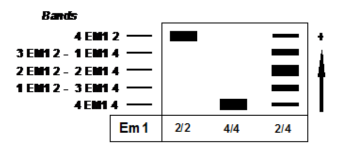
6. Recognition of the alleles encoding isoenzymes

6.1 Recognition of the alleles encoding ME

6.1.1 Genetic interpretation of the zymogrammes

Enzyme	Quaternary structure	Locus	Alleles
Malic enzyme (ME)	Tetrameric	Me1	2 4

6.1.2 Schematization of the zymogrammes

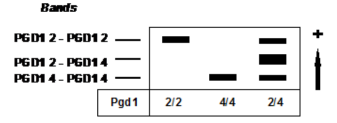


6.2 Recognition of the alleles encoding PGD

6.2.1 Genetic interpretation of the zymogrammes

Enzyme	Quaternary structure	Locus	Alleles
6-phosphogluconate dehydrogenase (PGD)	Dimeric	Pgd1	2 4

6.2.2 Schematization of the zymogrammes



Two migration zones can be observed; only the slowest migrating bands are polymorphic.

Common Sunflower, 2020-05-07

40

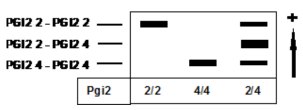
6.3 Recognition of the alleles encoding PGI

6.3.1 Genetic interpretation of the zymogrammes

Enzyme	Quaternary structure	Locus	Alleles
Phosphoglucoisomerase (PGI) Dimeric	Pgi2	2 4

6.3.2 Schematization of the zymogrammes

Bands



There are two migration zones; only the slowest migrating bands are scored.

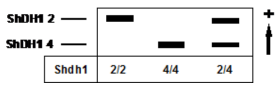
6.4 Recognition of the alleles encoding ShDH

6.4.1 Genetic interpretation of the zymogrammes

Enzyme	Quaternary structure	Locus	Alleles
Shikimate dehydrogenase (ShDH)	Monomeric	Shdh1	2 4

6.4.2 Schematization of the zymogrammes

Bands



6.5 Recognition of the alleles encoding PGM

6.5.1 Genetic interpretation of the zymogrammes

Enzyme	Quaternary structure	Locus	Alleles
Phosphoglucomutase (PGM)	Monomeric	D	2

Pgm4

6.5.2 Schematization of the zymogrammes

Bands PG 14 2 -PG M4 4, PG M3 4 · Pgm4 2/2 4/4 2/4

2
4

Common Sunflower, 2020-05-07

41

Several migration zones can be observed; only the fastest zone is polymorphic.

There is another gene which has not been considered. This has been designated Pgm3, encoding an enzyme which comigrates with PGM4 4.

So, the genotypes Pgm4 2/2 and Pgm4 2/4 give a two-band zymogramme. These both genotypes differ only by relative band intensities.

[End of Annex and of document]

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