# IN VITRO PROPAGATION OF SOME FRUIT SPECIES.

B- In vitro propagation of jojoba (Simmondsia chinensis Link, Schnider) plant.

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# **ABSTRACT**

Vegetative propagation of jojoba plant is difficult by traditional methods. A factorial experiment was conducted to develop a protocol for cloning jojoba through tissue culture technique. In this concern, shoot tips and nodal cuttings were prepared from jojoba plants. After sterilization, the explant were initiated on three culture media i.e. MS, B5 and WPM each at either full, one half or one fourth strength, these media supplemented with 0.1 mg/L IBA, 1.0 mg/L BA for establishment stage. After four weeks, MS medium gave the best results with the three measurements (survival %, shoot length and number of leaflets) full strength media proved to be the more suitable for the three measurements. Shoot tips surpassed nodal cutting explant. On the other hand, nodal cuttings which cultivated in quarter WPM had the lowest value in this respect. The newly formed shoots were transferred to the same media supplemented with either BA; Kinetin or 2ip at the concentration of 2, 4, 6 mg/L for each through proliferation stage. Full strength MS medium supplemented with 2 mg/L BA was superior and had the greatest number of shoots during the three subcultures. While the reverse was true with kinetin at 6 mg/L added to full strength WPM. Microshoots were rooted in the same half strength media with or without activated charcoal supplemented with 7 mg/L IBA + 1 mg/L NAA plus either 1 or 1.5 mg/L caffeic acid. The data revealed that WPM was most suitable for the three rooting growth measurements (rooting percentage, number of roots/plantlet and average root length) the presence of activated charcoal increased significantly the three rooting growth measurements iBA at (7 mg/L) + NAA at (1mg/L) + caffeic acid at (1 mg/L) gave the highest value of rooting measurements. While, the reverse was detected by the charcoal omitted B<sub>5</sub> medium supplemented with IBA at (7 mg/L) + NAA at (1 mg/L) + caffeic acid at (1.5 mg/L) during the two seasons of study. The plantlet produced from the best treatments of each medium, during the rooting stage were transplanted to (300 ml) plastic pots containing autoclaved transplanting media ( vermiculite : peat moss: sand mixed by volume (1:1:1) and maintained in green house for four weeks to investigate their effect on survival %, plant height and number of leaves per plant during acclimatization stage. The obtained results could be summarized as follows:rooted plantlet in 1/2 strength WPM + IBA (7 mg/L) + (1.0 mg/L) NAA + (1 mg/L) caffeic acid + 1.0 mg/L activated charcoal gave the highest value of rooting growth measurements while the reverse was true with rooted plantlet in 1/2 strength B5 + IBA (7.0 mg/L) + (1.0 mg/L) NAA + (1.0 mg/L) caffeic acid + 1.0 mg/L activated charcoal.

## INTRODUCTION

Jojoba plant (Simmondsia chinensis, Link, Schneider) which pronounced as ho-ho-ba belongs to family Simmondisceae. This plant is native to the arid zones of USA and Mexico.

Its natural distribution lies between 25 and 34 latitudes in an area, which closely approximates the Sonorant Desert (Gentry, 1958). Jojoba plant has currently received a special attention since its seeds contain liquid waxy

called jojoba oil. This oil is very similar to that obtained from sperm whale. The liquid wax of jojoba is used as a natural base for wide range of cosmetics and medicinal products, in addition, it has heat resistant lubricating properties and useful in chemical industry (Naqvi et al., 1988).

Clonal propagation exhibited elite individuals of known sexuality and, special relevance in order to make sure of the number of productive plants in a given plot. Its vegetative propagation is difficult by traditional methods (Yermanos, 1979). Furthermore, there are other horticultural limitations since, only a few cuttings can be obtained besides, the hardened terminal shoots are taken during a particular period of the year.

Several attempts have been made to develop tissue culture methods for propagation of *Simmondsia chinensis* (Aragao, 1977), but no success could be achieved in transferring the *in vitro* – regenerated plants to soil. The transplanting of *in vitro* rooted shoots of jojoba in recent pulplication is very scarce. We propose this study to develop a protocol for cloning *Simmondsia chinensis*, through tissue culture and successful transplantation of the *in vitro* – raised plants to soil.

#### MATERIALS AND METHODS

The present study was conducted in the Tissue Culture Laboratory, Horticulture Research Institute, Agriculture Res. Center during two seasons of 2002 and 2003. Generally, the following experiments were carried out:

I. Establishment stage:

In this stage, it was aimed to determine the suitable explant type (shoot tip & nodal cutting); kind of media (MS Murashige and Skoog, (1962); B5 Gamborg, et al., (1968) and WPM Lloyd and McCown (1980)) and media strength (full; half and quarter) by which more success could be achieved through the direct regeneration.

New growing shoots were taken at the beginning of the growing season (early March), washed with running water, and cut into either shoot tip or nodal cutting with about 10 mm length for each. Then explants were washed with tap water for one hour and soaking for 20 minutes in a commercial bleach "Clorox" (5.25 % sodium hypochlorite) at 20 % with two drops of tween-20, and then rinsed three times in sterilized distilled water for ten minutes per each to remove any residues of Clorox.

The prepared explants were cultured on three different nutrient media (MS,  $B_5$  or WP) each supplemented with 3 % sucrose; 0.1 mg/L IBA; 1.0 mg/L.BA (6- benzyl adenine) then solidified by using purified agar (Bacto-Difco agar) at 0.7 %. The PH of the media was adjusted to (5.6 to 5.8). Then, the media dispended into 100 ml glass jar each contained 25 ml medium then wrapped with plastic screw cap and sterilized. The media were autoclaved at (15 lb/in²) and 121°C for 20 minutes. All cultures were incubated under conditions of 25°c  $\pm$  2; 16 hours artificial light (fluorescent light at 30  $\mu$ M/ hz /sc) and 8 hours darkness.

The investigated treatments in this study which representative of the differential 18 combinations between 2 explant types (shoot tip and nodal cutting) x 3 media type (MS,  $B_5$  & WP) X 3 media strength (full,  $\frac{1}{2}$  and  $\frac{1}{4}$ ) were arranged in a factorial experiment using the complete randomized

design with three replications per each treatment .Every replicate was represented by 10 jars each contained 4 cultured explants.

After four weeks from culturing and incubation, data on survival % of cultured explants, shoot length and number of leaflets / shoot in response to investigated treatments (18 combinations) were recorded.

#### 2. Proliferation "shoot multiplication" stage:

Proliferated shoots throughout the previous stage i.e. establishment "  $1^{\underline{st}}$  stage " were used for the multiplication stage. Hence, regenerated shoots of both shoot tip and nodal cutting were collected and cultured preliminary on solid Murashege and Skoog (MS), Gamborg (B5) and Woody plant (WP) media supplemented with several growth regulators i.e., combinations of the cytokinin with auxin, (0.1mg/L) IBA, (30gm/L) sucrose and one of 3 cytokinin kinds i.e., kinetin; BA (benzyl adenine) or 2IP (isopentel adenin) at concentration of (2,4,6 mg/L) for each . Each medium (MS, B5 and WP) was supplemented with (100 mg/L) myo-inositol, 3 % sucrose, pH was adjusted at 0.7 %. Media were autoclaved at (1.5 kg / cm²) and 121°C for 20 min, then left to cool 24 hrs.

A factorial experiment using the complete randomized design with three replications was conducted for arranging the investigated 27 treatments i.e, various combinations between 3 media types X 3 cytokinin kinds X 3 concentrations of growth regulators (2, 4 and 6 mg) treatments. Every replicate was represented by five jars, each contained (40 ml) medium and 2 cultured explants. Data on the number of proliferated shootlets per each original one through 3 subcultures included in this stage were recorded.

# 3. Rooting stage:

proliferated shoots were taken and separated from each other under aseptic conditions and cultured on half-strength Murashege & Skoog (MS), Gamborg (B<sub>5</sub>) and Woody plant (WP) media supplemented with (30 g/L) sucrose and (7 g/L) purified Bacto - Difico agar with activated charcoal (1 g/L) or without. Rooting media were also varied pertaining auxin treatments i.e., IBA 7 mg + NAA 1 mg/L + Caffiec acid at either 1.0 or 1.5 mg/L, pH was adjusted at (5.6-5.8) and the media were autoclaved. Elongated shoots were transferred to jars containing (40 ml) of the abovementioned rooting media and incubated for one week in the dark and for 3 weeks in light. Where, rooting%; number of rootlets per plantlet and average length of each were recorded in response to the investigated treatments which were representative of 12 combinations between (3media types x 2 activated charcoal x 2 caffiec acid levels) with 3 replications.

## 4. Acclimatization stage:

Produced Jojoba plantlets were washed with tap water (Ebida, 1991 and Fassuliotis and Nelson, 1992) then dipped in Rhizolix solution (1.0 g/L) as fungicide for (10 min) prior to transplanting in (300ml) plastic pots containing autoclaved transplanting medium (vermiculite: peat moss: sand at (1:1:1) and maintained in green house for four weeks.

Pots were arranged then covered with polyethylene bags to maintain high relative humidity around the plants in green house (Fassuliotis and Nelson, 1992). After two weeks, the polyethylene bags were partially removed to allow air circulation (Ali et al., 1990), and later removed after

other two weeks (Smith, 1981). Plantlets were irrigated with half strength (MS,  $B_5$  and WP) maintenance medium (free hormone medium) during the period of hardining (Ebida, 1991). The irrigation was applied depending on the requirement of plantlets. Pests and disease control program was followed as recommended.

Data were recorded after one month of transplanting as follow:

- 1- Survival percentage.
- 2- Plant length (cm).
- 3- Number of leaves / plant.

#### Statistical analysis:

Data obtained were statistically analysed according to (Snedecor and Cochran, 1980) and significant differences among means were determined by Duncan's multiple range test (Duncan, 1955).

## RESULTS AND DISCUSSION

## 1-Establishment stage:

## 1-1 Survival percentage:

Concerning the specific effect of different factors involved in this study i.e., explant type, media strength and media type on survival percentage data presented in Table (1) showed that shoot tip recorded higher value of survival percentage than nodal cuttings during both seasons of study.

As for the specific effect of media strength, Table (1) reveals that, full strength media was superior as it exhibited statistically the highest value of survival %, followed in a descending order by half strength media which ranked the second, while quarter strength media ranked last to represent the inferior strength during 2002 and 2003 seasons.

Regarding the specific effect of media type, data obtained revealed, the superiority of (MS) medium over the other ones which showed higher survival % Moreover, (B<sub>5</sub>) medium ranked statistically  $2^{\underline{nd}}$  while (WP) medium ranked  $3^{\underline{nd}}$  during  $1^{\underline{st}}$  and  $2^{\underline{nd}}$  seasons.

Concerning the interaction effect of various combinations on survival %, data obtained revealed that combinations representing cultured shoot tips or nodal cutting on full strength of 3 media (especially MS) and to great exetent shoot tip on half MS strength exhibited statistically the highest survival % during two seasons of study.

On the contrary, cultured explants (shoot tip & nodal cutting) on one fourth WP medium exhibited the least survival % during both seasons of study.

Moreover, other combinations were in between the above-mentioned two extremes. These results go in line with Turk et al., (1992); Zaman et al., (1998) and Silva et al., (2003).

Specific and interaction effects of explant type, media strength, media type and their combinjojoba Simmondsia chinensis during establishment stage ( 2002 & 2003 ations on survival % of Table (1):

Explant type Media type B5 Shoot tip MS	0		Strength of media	edia		Stre	Strength of media	edia	
Shoot tip	type	Full	Half	Quarter	Mean*	Full	Half	Quarter	Mean*
Shoot tip			2002				0 11 11	2003	
Shoot tip	B5	82.67ab 78.00c	78.00c	71.00f		82.33ab	82.33ab 77.98cd	68.67f	
	MS	83.20a	83.20a 81.67ab	78.20c	77.38A	83.00a	81.60ab	76.67d	76.86A
	WP	81.65ab	73.67e	66.33g		82.25ab	82.25ab 73.60e	66.67g	
	B5	82.30ab 77.00cd	77.00cd	70.00f		82.00ab	76.67d	69.00f	
Nodal cutting	MS	82.35ab	81.50b	76.00d	76.76B	81.33ab	81.33ab 79.00c	75.00e	76.16B
in the second	WP	81.20b	74.20e	66.30g		82.50ab	82.50ab 74.00e	66.30g	
Mean **		82.24A	77.67B	71.31C		82.01A	77.14B 74.89C	74.89C	i de Ge
Mean ***		85	MS	WP		B5	MS	WP	
		76.83B	80.49A	73.89C		76.11B	76.11B 79.63A 73.80C	73.80C	

\*, \*, \*\*\* Refer to specific effect of explant type, media strength and media type treatments, respectively. Capital and small letter / swere used for distinguishing between values of specific and interaction effects, respectively whereas means followed by the same letter/s were not significantly different at 5 % level.

## 2- Length and number of leaves per jojoba explant:

In this regard specific effect of three studied factors i.e., explant type (shoot tip & nodal cutting), media strength (full, half and quarter) and media type ( $B_5$ , MS and WP); as well as their possible combinations were investigated pertaining the response of average shoot length and number of leaflets per each.

Referring the specific effect of explant type, it is quite clear as shown from Table (2), that shoot tip had higher values of both shoot length and number of leaflets/shoot than nodal cutting during the two seasons of study.

As for the specific effect of media strength, data displayed that full strength induced statistically the tallest shoot with highest number of leaflets/shoot followed in descending order by half strength and quarter strength, whereas differences were significant during the 2002 and 2003 experimental seasons .

With regard to the specific effect of media type, the results show that (MS) medium proved to be the best medium in establishment stage which exhibited the highest values of both shoot length (cm.) and number of leaflets/shoot while (WP) medium was the least effective during the two seasons of study.

Referring the interaction effect: Table (2) and photo (1) & (2) show, an obvious variances between combinations of explant type; media strength and media type.

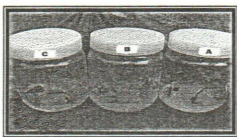


Photo (1)

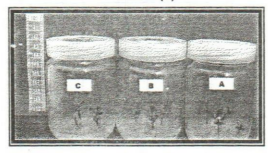


Photo. (2)

Photos. (1 & 2): Effect of explant type, media type and media strength on measurements of establishment stage of jojoba Simmondsia chinensis explant:

Photo 1- Shoot tip in full strength media (A: MS, B: B<sub>5</sub>, C: WP) Photo 2- Nodal cutting in full strength media (A: MS, B: B<sub>5</sub>, C: WP)

Table (2) :Specific and interaction effects of explant type, media strength, media type and their combinations on shoot length (cm.) and No. of leaflets / shoot of jojoba Simmondsia chinensis during establishment etada (2002 & 2003 seasons)

	Media strength	Sh	Shoot length (cm.)	(cm.)		No. o	No. of leaflets/shoot	hoot	
2	Media type	Full	Half	Quarter	Mean*	Full	Half	Quarter	Mean*
				2002					
-	B5	2.08b	1.78c	1.45fg		5.00c	4.50f	3.75i	
-	MS	2.45a	2.01b	1.80c	1.81A	5.75a	4.58ef	4.00h	4.48A
-	WP	1.81c	1.58de	1.30hi		4.73de	4.45f	3.58ij	
-	B5	1.96b	1.73cd	1.39gh		4.83cd	4.25g	3.50	
nodal cutting	MS	2.34a	2.02b	1.65d	1.72B	5.33b	4.75de	3.75	4.31B
	WP	1.68cd	1.52ef	1.23i		4.58ef	4.41fg	3.41j	71
Mean **	** U	2.05A	1.77B	1.47C		5.04A	4.49B	3.67C	
	400	85	MS	WP		B5	MS	WP	
Mean		1.738	2.05A	1.52C	100	4.31B	4.69A	4.19C	
			100	2003			63	5 65 17	
	B5	2.07b	1.77ef	1.40j		5.20b	4.65bc	3.8ef	
_	MS	2.43a	2.02bc	1.77ef	1.79A	5.92a	4.70bc	4.15d-f	4.57A
_	WP	1.80de	1.57hi	1.25k		4.6bc	4.30cd	3.75ef	
	B5	1.92cd	1.72e-g	1.38j		4.67bc	4.33cd	3.58f	
Nodal cutting	MS	2.32a	1.96bc	1.60g-l	1.70B	5.76a	4.42cd	3.83ef	4.28B
_	WP	1.65f-h	1.50ij	1.20k		4.30cd	4.08de	3.55f	
Mean **	n **	2.03A	1.76B	1.43C		5.07A	4.41B	3.78C	2 7
	****	85	MS	WP		B5	MS	WP	
Mean		1.718	2.02A	1.49C		4.37B	4.80A	4.10C	

\*\*\*\*\*\* Refer to specific effect of explant type, media strength and media type , respectively. Capital and small letter / s were used for whereas means followed by the same letter/s were distinguishing between values of specific and interaction effects, respectively not significantly different at 5 % level.

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Whereas, the full strength (MS) medium was more effective and showed the highest value of shoot length and greatest number of leaflets/shoot. Conversely, both shoot tip and nodal cutting which cultured on (WP) quarter medium strength gave shortest shoots with lowest number of leaflets/shoot followed in an increasing order by both shoot tip and nodal cutting which cultured on ( $B_5$ ) quarter medium strength ranked second for number of leaflets/shoot during 2002 and 2003 experimental seasons.

The obtained results go in line with the findings of Tabachnik & Kester

(1977), Hammerschlage (1982) and Saker et al., (1999).

## 2- Multiplication stage:

In this respect specific effect of three studied factors i.e., media type (B $_5$ , MS & WP); cytokinins kind (BA; 2IP and kinetin) and applied concentrations (2, 4, 6 mg/L) of these three cytokinins, as well as their possible combinations were investigated pertaining the response of number of proliferated shoots. Data obtained four weeks after  $1^{st}$ ,  $2^{nd}$  and  $3^{rd}$  subcultures of multiplication stage are presented in Table (3).

Concerning the specific effect of media type, it is quite clear as shown from Table (3) that, Murashige & Skoog (MS) medium was the superior through three subcultures where the greatest (number of developed shootlets) was resulted followed in a descending order by Gamborg (B<sub>5</sub>)

medium and Woody plant (WP) medium during two seasons.

With regard to the specific effect of concentration of added cytokinins, Table (3) shows that, 2 mg/L resulted significantly in the greatest number of developed shoots descendingly followed by 4 mg/L and 6 mg/L. Differences were significant during both 2002 and 2003 experimental seasons.

Referring the specific effect of cytokinins kind it is so clear to be noticed that BA (benzyl adenine) exhibited significantly the highest value for number

of proliferated shoots descendingly followed by 2iP (isopentel adenine) while Kinetin ranked the last. Differences were significant during both seasons.

Concerning the interaction effect: Data obtained during both 2002 and 2003 experimental seasons as shown from Table (3) and photo (3) displayed that the highest number of developed shoots were significantly in close relationship with Murashige & Skoog (MS) medium supplemented with BA at (2 mg/L). On the contrary, the least values for the number of developed shoots where coupled with Woody plant (WP) medium supplemented with Kinetin at (6 mg/L) during the study. However, other combinations were in between the previously mentioned two extremes during three subcultures. These results are in general agreement with those found by Liorente et al., (1998), Chitra & Pandmaja (1999), Join & Bubbar (2000) and Erig & Schuch (2003).

Specific and interaction effects of media type; growth regulators( kind & concentration) and their combinations on number of shoots (4 weeks later ) during multiplication stage of jojoba (Simmondsia Table (3):

٦.	1	s Means'	9			2.58l 3.70B	1.67p		2.92j 4.44A	1.83op	_	2.17mn 3.29C	1.95no	2.41C	Kinetin	2.56C			2.67k-m 3.66B	1.750	_	2.91j-I 4.62A	Ono	_	2.30mn 3.313C	)Bno	2.48C	Kinotin
Thrid Sub.	Concentrations of	growth regulators	<u>.</u>			+	-	-	-									3f 3.0	gh 2.6		-	3e 2.9				-		
Th	centra	vth reg	4		4.67f	4.00g	2.66kl	5.67d	5.00e	2.83jk	3.67h	3.33	2.75]-	3.85B	2iP	4.11B	1887	4.66f	3.67gh	2.66k-m	5.83d	5.33e			3.50g-l	2.70	3.90B	0
	Con	grov	2		7.00c	4.64f	3.16i	7.67a	7.33b	3.30	5.60d	5.00e	2.87	5.17A	BA	4.76A		6.67c	4.60f	3.29h-j	8.33a	7.66b	3.50g-l	5.50de	4.55f	2.88-1	5.22A	40
		Means*				3.30B			4.09A			2.95C							3.241B	100	N	4.037A			2.972C			
d Sub	1s of	ators	9		2.29kl	2.16lm	1.50n	2.75	2.50jk	1.67n	2.09m	2.00m	1.48n	2.05C	Kinetin	2.11C	THE WALL	2.32k-m	2.15I-n	1.420	2.67jk	5.10ef 2.33k-m	1.66no	1.92m-o	2.12l-n	1.420	2.00C	
Secound Sub	Concentrations of	growth regulators	4		4.67f	3.33h	2.10m	5.67d	5.00e	2.31kl	3.30h	3.28h	2.08m	3.53B	2iP	3.77B	9 30KT 0	4.00hi	3.67i	2.17I-n	5.66cd	5.10ef	2.30k-m	3.67i	3.00	2.25k-n	3.53B	2
	Conc	grow	2	2002	6.30c	4.66f	2.70i	7.67a	6.67b	2.60ij	5.45d	4.33g	2.55ij	4.77A	BA	4.46A	2003	6.00bc	4.60fg	2.83jk	7.66a	6.33b	2.66j-l	5.33de	4.30gh	2.65j-1	4.71A	-
		Means*		2(		2.92B			3.93A			2.48C					2		3.00B			3.88A			2.600C			
Cirk	Sub.	itors	9		2.16ik	2.00k	1.42mn	2.50	2.25	1.58lm	1.98k	1.72	1.25n	1.87C	Kinetin	1.98C		2.33f-h	2.15g-I	1.33	2.42fg	2.20q-i	1.58	1.83h-i	1.71	1.33	1.87C	
Evict Cub	LIST	concentrations of growth regulators	4		3.669	3.659	2.10k	5.67c	4.33e	2.17Kl	3.00h	2.60	1.691	3.21B	2iP	3.44B		3.55e	3.50e	2.18a-l	5.17c	4.50d	2.10g-I	3.67e	2.75f	1.67	3.23B	
Evice Su	0 000	Conc	2		4 67 d	4.17ef	2.48i	7.66a	6.67b	2.58i	4.00f	3.60g	2.45i	4.25A	BA	3.92A		5.00c	4.31d	2.66fa	7.67a	6.66b	2.67fg	4.30d	3.62e	2.58fh	4.39A	
Minches of phones	Number or shoots	Growth	regulators type	- If a sample	BA	2iP	Kinetin	ВА	ZiP	Kinetin	BA	2iP	Kinetin	Mean**		Mean***		BA	ZiP	Kinetin	BA	2iP	Kinetin	BA	2iP	Kinetin	Mean**	
	:	Media	ry po			85	1		SM	)		WP							85			SM			WP			

Table (4): Specific and interaction effects of media type; activated charcoal; auxin treatments added to one half strength rooting medium and their combinations on rooting percentage; number of rootlets and average root length (cm.) through rooting stage of jojoba Simmondsia chinensis during (2002 & 2003 seasons).

Charcoal         Rooting percentage         A.Ch.         Mean**         Me		Treatments				4	Parameters	ers			
Number of roots   Auxins   A	:										Root length
Mithout   Mean**   Mithout   Mean*   Mean*   Mean**   Mithout   Mean**   Mithout   Mithout   Mean**   Mean**   Mithout   Mithout   Mean**   Mithout   Mithout   Mean**   Mithout   Mithout   Mean**   Mithout   Mithou	Media			Rootii	ng perce	entage		Nun	lber of	coots	(cm.)
With out   Without   Wit	type	Auxins	A.C	Sh.	RADONA	Mosn**	A	.ch.	Moon*	Moon**	A.Ch.
BA7ML + NAA 1ML + caffeic acid 1 ML   16.00e   17.33r   16.92C   1ML + NAA 1ML + caffeic acid 1.5 ML   16.00e   17.33r   16.92C   1ML + caffeic acid 1.5 ML   16.00e   17.00fg   16.00e   16.00e   16.00e   16.00e   16.00e   16.00e   17.00fg   16.00e   16				Without	Meall	Mean	With	Without	Mean	Mean	With
IBA7ML + NAA 1ML + caffeic acid 1 ML   18.00e   17.33r   16.92C   1ML + caffeic acid 1.5 ML + caffeic acid 1					2002						
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   1667g   15.67h   16.92C   1ML + caffeic acid 1.5 ML + caffeic acid 1.5 ML   16.33e   19.33d   18.73e   19.33d   18.73e   19.33d   18.73e   19.50d   1		IBA7ML + NAA 1ML + caffeic acid 1 ML	18.00e	17.33r		IBA7ML + NAA	3.15e	2.67f		IBA7ML + NAA	10.67cd
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   18.33e   19.33d   18.75B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   19.53d   19.50d   19.50d   19.50d   19.50d   19.50d   19.50d   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   19.53d   19.50d   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   19.30e   19.15B	82	IBA7ML + NAA 1ML + caffeic acid 1.5 ML	16.67g	15.67h	16.92C	1ML + caffeic acid 1 ML	2.83f	2.58f	2.81C	1ML + caffeic acid 1 ML	8.67ef
IBA7ML + NAA 1ML + caffeic acid 1.5 ML	000	IBA7ML + NAA 1ML + caffeic acid 1 ML	20.33e	19.33d	40 700	20.31A	3.67d	3.29e	2000	3.03A	11.00c
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   24.20a   22.67b   21.44A   15ML   19.53d   19.50d   21.44A   17.17B   4.00c   3.42e   3.18B   3.56A   3.18B   3.15B   3.85A   3.00d   19.50d   19.50d	MS	IBA7ML + NAA 1ML + caffeic acid 1.5 ML	18.33e	17.00fg	16.755	IBA7ML + NAA	3.28e	2.80f	3.200	IBA7ML + NAA	9.67c
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   19.53d   19.50d   18.0d   19.50d   19.50d	WPM	IBA7ML + NAA 1ML + caffeic acid 1 ML	24.20a	22.67b	21.44A	1ML + caffeic acid 1.5ML	4.67a	4.30b	4.10A	1ML + caffeic acid 1 .5ML	12.00a
BA7ML + NAA 1ML + caffeic acid 1.5 ML   17.30c   17.20h   19.30c   17.30c   17.30c   17.30c   17.30c   17.30c   18.53b   19.15b   19.00c   19.30c   19.00c   19.00c		IBA7ML + NAA 1ML + caffeic acid 1.5 ML	19.53d	19.50d		17.178	4.00c	3.42e		3.15B	9.31d
BA7ML + NAA 1ML + caffeic acid 1 ML		Mean**		18.53B			3.60A	3.18B			10.22A
IBA7ML + NAA 1ML + caffeic acid 1 ML   21.33c   17.20h   17.95C   1ML + caffeic acid 1.5 ML   17.67g   15.60i   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   18.00f   17.30gh   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   22.00b   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   24.00a   22.33b   21.48A   15.6ML   18.00b   19.60de   19					2003						
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   17.67g   15.60i   17.95C   1ML + caffeic acid 2.55f   2.33g   2.65C   1ML + caffeic acid 1.5 ML   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   22.03b   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   24.00a   22.33b   21.48A   1.5ML   18.03B   3.85A   3.04B   3.04B   20.00d   19.60de   18.00b   19.60de   18.00b   19.60de   18.00b   19.60de   19.50b   19.50b   19.50b   19.50b   19.50b   19.50b   19.50b   19.50b   19.60de   1		IBA7ML + NAA 1ML + caffeic acid 1 ML	21.33c	17.20h		IBA7ML + NAA	3.10e	2.60f		IBA7ML + NAA	10.60c
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   22.00b   19.30e   19.15B   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   24.00a   22.33b   18.00f   19.60de   18.00d   19.60de   18.00d   19.60de   18.00d   19.60de   18.00d   19.60de   19.00d   19.00d   19.60de   19.00d   19.00d	B2	IBA7ML + NAA 1ML + caffeic acid 1.5 ML	17.67g	15.60i	17.95C	1ML + caffeic acid 1 ML	2.55f	2.33g	2.65C	1ML + caffeic acid 1 ML	8.60ef
IBA7ML + NAA 1ML + caffeic acid 1.5 ML   18.00f   17.30gh   19.13b   IBA7ML + NAA 1ML + caffeic acid 1.5 ML   24.00a   22.33b   15.00c   19.60de   19.60de   18.03b   18.03b   3.85A   3.04B   2.82B   18.03b   3.38A   3.04B   2.82B   2.82B   2.050A   18.56B   2.050A   18.56B   2.050A   18.56B   3.38A   3.04B   3.04B	074	IBA7ML + NAA 1ML + caffeic acid 1 ML	22.00b	19.30e	0.77.07	21.03A	3.40d	3.33d	0 4 40	3.55A	11.10b
IBA7ML + NAA 1ML + caffeic acid 1.5 ML + c	MS	IBA7ML + NAA 1ML + caffeic acid 1.5 ML	18.00f	17.30gh	19.130	IBA7ML + NAA	3.17e	2.66f	0.140	IBA7ML + NAA	9.66d
IBA7ML + NAA 1ML + caffeic acid 1.5 ML         20 00d         19.60de         18.03B         3.58c         2.99f         2.82B           Mean***         20.50A         18.56B         3.38A         3.04B         2.82B	WPM	IBA7ML + NAA 1ML + caffeic acid 1 ML	24.00a	22.33b	21.48A	1ML + caffeic acid 1.5ML	4.50a	4.33b	3.85A	1ML + caffeic acid 1 .5ML	12.20a
20.50A 18.56B 3.38A 3.04B		IBA7ML + NAA 1ML + caffeic acid 1.5 ML	20.00d	19.60de		18.03B	3.58c	2.99f		2.82B	9.30cd
		Mean***	20.50A	18.56B			3.38A	3.04B			10.24A

small letter/s were used for distinguishing between values of specific and interaction effects,respectively whereas means ","and"" Refer to specific effect of media type; auxin treatment and activated charcoal added to rooting media, respectively. Capital and followed by the same letter/s were not significantly different at 5 % level.

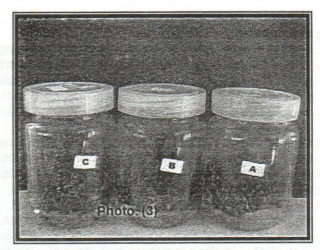


Photo. (3): Effect of cytokinins added at 3 levels (2, 4 and 6 mg/L) to three media through the 3<sup>rd</sup> subculture within multiplication stage of jojoba (Simmondsia chinensis).

A: Cultured explant in MS medium supplemented with (2mg /L) BA

B: Cultured explant in B5 medium supplemented with (2mg /L) BA

C: Cultured explant in WP medium supplemented with (2mg /L) BA

#### 3- Rooting stage:

In this regard, adding auxins IBA (7 mg/L); NAA at 1 mg/L and Caffeic acid at (1 or 1.5 mg/L) to half strength B<sub>5</sub>, MS & WP media either supplemented with (1.0 g/L) activated charcoal or not in combination were investigated after incubation for 4 weeks through rooting stage (either dark was applied at the 1<sup>st</sup> week or not ) regarding the influence on rooting percentage, number of developed rootlets per plantlet and average root length (cm.) of Jojoba plant. Data obtained are presented in Table (4) and Photo. (4).

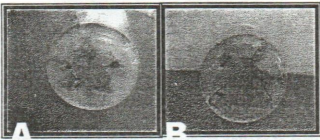


Photo. (4)

Photo. (4): Effect of combinations between media type, auxin treatments and charcoal adding on some measurements during rooting stage of jojoba (Simmondsia chinensis)

A: Cultured shootlets in 1/2 strength WP + IBA (7.0mg / L) + (1.0mg /L) NAA + (1.0mg /L) Caffeic acid without activated charcoal.

B: Cultured shootlets in 1/2 strength MS + IBA (7.0mg / L) + (1.0mg /L) NAA + (1.0mg /L) Caffeic acid without activated charcoal.

## 3. Rooting percentage:

Concerning the specific effect of media type on the rooting percentage, data showed that, Woody plant (WP) medium exhibited statistically the greatest rooting %, followed in a descending order by Murashige & Skoog (MS) medium and Gamborg (B<sub>5</sub>) medium which ranked last. Differences during both seasons were significant as the three media were compared each other.

Regarding the specific effect of adding activated charcoal to half strength media, data displayed that adding activated charcoal to rooting medium was effective. However, the activated charcoal omission reduced rooting percentage of jojoba plantlet during  $1^{\underline{st}}$  and  $2^{\underline{nd}}$  seasons.

As for the specific effect of two auxin treatments (7 mg/L IBA + 1 mg/L NAA +1.0 mg/L Caffeic acid) and (7 mg/L IBA + 1 mg/L NAA +1.5 mg/L Caffeic acid) added to half strength rooting media (supplemented with charcoal or not). Data obtained displayed that the auxin treatment with the lower Caffeic acid level (1 mg/L) was more suitable than the higher Caffeic acid rate (1.5 mg/L) during the two seasons of study.

Concerning the interaction effect, it could be safely concluded that half strength (WP) rooting medium supplemented with activated charcoal (1.0 g/L) plus IBA (7 mg/L) + NAA at (1 mg/L) + Caffeic acid at (1 mg/L) gained statistically the highest rooting % when subjected to darkness through  $1^{\underline{s}\underline{t}}$  week of incubation during the two seasons of study. Moreover, incubation of jojoba plantlets in half strength charcoal omitted (B<sub>5</sub>) medium supplemented with IBA at (7 mg/L) + NAA at (1 mg/L) + Caffeic acid at (1.5 mg/L) had the lowest value of rooting % during 2002 and 2003 seasons. In addition, other combinations were in between.

These results are in general agreement with the findings of, Magyar et al., (2001); Thomas, (2003) and Soliman (2004).

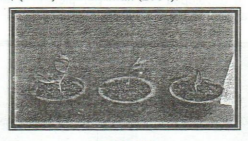


Photo (5)

- Photo (5): Effect of rooting media type, auxin treatments and activated charcoal during rooting stage on some measurements of jojoba (Simmondsia chinensis) after acclimatization stage:
  - A: Rooted plantlets on 1/2 strength WP + IBA (7.0 mg / L) + (1.0 mg /L) NAA + (1.0 mg /L) Caffeic acid + (1.0g/L) activated charcoal
  - B: Rooted plantlets on 1/2 strength MS + IBA (7.0 mg / L) + (1.0 mg /L) NAA + (1.0 mg /L) Caffeic acid + (1.0g/L) activated charcoal
  - C: Rooted plantlets on 1/2 strength B5 + IBA (7.0 mg / L) + (1.0 mg /L) NAA + (1.0 mg /L) Caffeic acid + (1.0g/L) activated charcoal

## 3.2- Number of roots per plantlet and root length (cm):

Regarding the response of number of the developed rootlets and their average length to the specific effect of investigated factor i.e, media type; auxin treatments and activated charcoal added through rooting stage, Table (5) displays that the greatest number of roots per plantlet and tallest roots were detected by rooted shortest on (WP) medium followed in a descending order by those on (MS) medium while (B5) medium had statistically the lowest values in this concern during two seasons of study.

As for the influence of adding activated charcoal at (1 gm/L) to half strength medium, data revealed that, the number of roots per plantlet and root length were significantly depressed on the charcoal omitted as compared to analogous one supplemented with charcoal.

As for the specific effect of auxin treatments; data obtained displayed that IBA at (7 mg/L) + NAA at (1 mg/L) + (1 mg/L), Caffeic acid treatment significantly increased the number of roots/plantlet and root length as compared with IBA at 7 mg/L + NAA at 1 mg/L + Caffeic acid at 1.5 mg/L during 2002 & 2003 seasons.

Concerning the interaction effect of various combinations between media type x auxin treatments x charcoal added), Table (5) and photo (4) show that half strength (WP) medium supplemented with (7 mg/L) IBA + (1 mg/L) NAA + (1 mg/L), Caffeic acid and activated charcoal gave significantly the greatest number and tallest rootlets per plantlet. On the contrary, adding IBA at (7 mg/L) + NAA at (1 mg/L) + Caffeic acid at (1.5 mg/L) without activated charcoal to half strength (B<sub>5</sub>) medium gave the lowest number of rootlets/plantlet and shortest rootlets during the two seasons of study.

In addition, other combinations were in between as compared to the previously mentioned two extents.

These results are in general agreement with the findings previously mentioned by Ishida et al., (1989); Vasar et al., (2000) and Soliman (2004). However, the presence of activated charcoal in rooting medium was in general agreement with the findings of Bondok et al., (1989); Fouad et al., (1995) and Soliman (2004).

# 4 - Acclimatization stage:

In this stage:- The plantlets produced from the best treatments (rooting media X auxins and charcoal added) through the previous stage (rooting) were chosen and cultivated on transplanting medium consisting of (vermiculite: peat moss: sand) at (1:1:1) ratio by volume for acclimatization stage.

Table (5) shows the effect of some Specific treatments used in rooting stage on survival% and some growth parameters (shoot length and number of leaves) during acclimatization stage.

Rooted plantlets in half strength WPM + IBA at 7mg/L+ NAA at 1mg/L+ Caffiec acid at 1.0 mg/L + 1.0 mg/L activated charcoal gave the highest survival%, tallest shoots and higher number of leaves followed in descending order by rooted plantlets in half strength MS medium + IBA at 7mg/L+ NAA at 1mg/L+ Caffiec acid at 1.0 mg/L + 1.0 mg/L activated charcoal. While rooted plantlets in half strength B5 medium + IBA at 7mg/L+ NAA at 1 mg/L+ Caffiec

acid at 1.0 mg/L + 1.0 g/L activated charcoal had the least values during the two seasons of study.

These results are in general agreement with the finding of Hoffmann et al., (1999); Benzioni et al., (2003) and Soliman (2004).

Table (5): Comparison between the most effective three rooting treatments (rooting medium X auxins and charcoal added) on survival %; shoot length (cm) and number of leaves of acclimatized newly regenerated jojoba plantlets during 2002 and 2003 seasons.

Parameters	Sur	vival	Shoot	length	No. I	eaves
Treatments	2002	2003	2002	2003	2002	2003
½ strength WP +IBA(7 mg/L) + NAA (1 mg/L) + Caffeic acid (1 mg/L) + A. C.1g/L		76.00 a	10.00 a	10.20 a	11.00 a	11.20 a
$\frac{1}{2}$ strength MS +IBA(7 mg/L) + NAA (1 mg/L) + Caffeic acid (1 mg/L) + A. C.1g/L		74.30 b	9.20 b	9.00 b	10.00 b	10.25 b
$\frac{1}{2}$ strength B5 +IBA(7 mg/L) + NAA (1 mg/L) + Caffeic acid (1 mg/L) + A. C.1g/L		71.60 c	7.67 c	7.66 c	9.20 c	8.83 c

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الإكثار بتقنية زراعة الأنسجة لبعض أنواع الفاكهة: ب- إكثار الهوهوبا بتقنية زراعة الأنسجة محمد عبدالوهاب خميس - وفاء توفيق سعيد - أحمد حسن جاد الحق ١ كلية الزراعة - جامعة بنها. ٢ - قسم بحوث الزيتون والمناطق شبه الجافة - معهد بحوث البساتين - مركز البحوث الزراعية - الجيزة - مصر.

أجريت هذه الدراسة في موسمي ٢٠٠٢ ، ٢٠٠٣ بمعمل زراعة الأنسجة بمعهد بحوث البساتين على نبات الهوهوبا بهدف تطوير اكثارها بطريقة زراعة الأنسجة لصعوبة اكثارها بالطرق التقليدية ويستلزم ان تجهز العقل في فترة معينة ومحدودة بالإضافة إلى ذلك فإن نسبة نجاحها قليلة وعلى ذلك فقد تم تجريب عدة معاملات خلال المراحل التالية (الأساس التضاعف التجذير والاقلمة) وقد تم دراستها كالأتي: ولا: مرحلة الأساس:

فى هذه المرحلة اجريت تجربة عاملية لدراسة التأثير النوعى لكل من المنفصل النباتى ( البرعم الطرفى والعقلة ذات البرعم الواحد ) وكذلك نوع البيئة (B5, MS, WPM) وتركيز املاحها الأساسية (كاملة ونصف وربع تركيز على نسبة البقاء ومتوسط طول الفريخات وعدد الوريقات المتكونة عليها) فبعد إجراء التعقيم للمنفصلات النباتية ثم زراعتها على البيئات الغذائية السابقة الذكر مضاف إلى كل منها ١٠، ملجم/ لتر الدول حامض البيوتيريك ١ ملجم/ لتر بنزيل ادنين وبعد أربعة أسابيع من الزراعة أظهرت النتائج المتحصل عليها الأتى:

■ تفوق البرعم الطرفى على العقلة ذات البرعم الواحد في تسجيل أعلى نسبة بقاء وكذلك طول الفريخات وعدد الأوراق المتكونة عليها.

■ أثبتت بيئة موراشيج وسكوج كاملة القوة أنها أفضل البيئات في تسجيل أعلى القيم للقياسات الثلاثة السابقة الذكر وكان العكس صحيحا مع العقلة ذات البرعم الواحد والتي تم زراعتها على بيئة الأشجار الخشبية ذات الربع تركيز .

ثاتيا: مرحلة التضاعف:

فى هذه المرحلة تم إعادة الزراعة الفريخات الناتجة من مرحلة الأساس على نفس البينات الغذائية السابقة الذكر ذات القوة الكاملة المضاف إليها ثلاثة أنواع من السيتوكينينات ( بنزيل أدنين وأيزوبنتيل أدنين وكينيتين ) كل بثلاث تركيزات هى ( ٢٠٤،٢ ملجم / لتر) فى تباديل وتراكيب مختلفة بينها لدراسة تأثيرها على عدد الأفرخ المتكونة وقد أوضحت الدراسة النتائج التالية :

 تفوقت بيئة موراشيج وسكوج (MS) في زيادة عدد الفريخات الحديثة المتكونة على بيئتي جامبورج وبيئة الأشجار الخشبية .

البنزیل أدنین بترکیز ۲ ملجم/لتر کان أکثر تفوقا فی هذا الشأن .

إضافة الكينيتين بتركيز ٦ ملجم/ لتر على بيئتى جامبورج أو الأشجار الخشبية أعطى أقل عدد تفرعات وهذا خلال الثلاث نقلات (subculture).

ثالثًا: مرحلة التجذير:

تم تجذير الأفرخ الجديدة المتكونة في مرحلة التضاعف على البينات الثلاثة السابقة الذكر ذات النصف تركيز لأملاحها والمحتوية على الفحم النشط بتركيز ١ جم / لتر أو الخالية من الفحم النشط والمضاف اليها ٧ ملجم / لتر اندول حامض البيوتيريك (١هم / لتر) + نفتالين حامض الخليك بتركيز (١ أو ١,٥ مجم / لتر) .

فقد درس تاثير التفاعل للتراكيب المختلفة على إستجابة قياسات التجنير الثلاثة (نسبة التجذير عدد الجذور المتكونة \_ متوسط طول الجذور) وقد أوضعت الدراسة النتائج التالية : \_

■ سجلت بيئة الأشجار الخشبية (WPM) أعلى قيمها لنسبة التجذير وعدد الجذور المتكونة ومتوسط طول الجذر.

إضافة الفحم النشط (١جم/لتر)إلى البيئات أدى إلى زيادة معنوية للقياسات الثلاثة خلال موسمى الدراسة .

■ إضافة ٧ مجم/لتر إندول حامض البيوتيريك + ١ ملجم / لتر نقثالين حامض الخليك + ١ مجم / لتر حامض الخليك + ١ مجم / لتر حامض الكافييك أعطى أعلى قيم لمقاييس التجذير الثلاثة السابقة الذكر، وكان العكس صحيحا مع بيئة جامبورج (B5) الخالية من الفحم النشط والمضاف إليها ٧ ملجم/ لتر إندول بيوتيريك (B5) + ١ ملجم/ لتر يقالين حامض الخليك (NAA) + ١,٥ ملجم/ لتر حامض الكافييك خلال موسمى الدراسة

رابعا: مرحلة الأقلمة:

في هذه المرحلة اجريت تحت ظروف الصوبة الزجاجية حيث تم نقل نباتات الهوهوبا الناتجة من افضل معاملة لكل بيئة من البيئات الثلاثة (B5, WPM, MS) المستخدمة تحت الدراسة في مرحلة التجذير الأقلمتها وذلك بغسلها بماء الصنبور وغمسها في محلول ريزولكس ثم تفريدها في أصحص بالستيك (٣٠٠ مم) مملوءة بمخلوط معقم من البيت موس والفيرميكوليت والرمل بنسبة حجميه (١:١:١) لمدة ٤ أسابيع لدراسة نسبة البقاء وطول النبات وعدد الأوراق لكل منها وقد أوضحت النتائج المتحصل عليها:

■ النباتات المجدرة على بيئة الأشجار نصف تركيز والمحتوية على ٧ ملجم / لتر إندول حامض البيوتيريك
 + ١ مجم / لتر نفثالين حامض الخليك + ١ مجم / لتر حامض الكافييك + ١ جم فحم نشط كانت الاكثر
 تفوقا بالنسبة لمقاييس الاقلمة الثلاثة (نسبة البقاء \_\_\_ طول النبات وعدد الأوراق) .

■ النباتات المجدرة على بيئة B<sub>5</sub> نصف تركيز والمضاف اليها ٧ مجم /لتر اندول حامض البيوتيريك(IBA) + 1 ملجم نفتالين حامض الخليك (NAA) + 1 ملجم حامض الكافييك + 1 جم فحم نشط أظهرت أقل قيم في هذا الصدد .