Germination in Vitro of Brassicaceae (*Sinapis arvensis* L.) in the Northern Region of Tlemcen (Algeria)

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Abstract: This ecophysiological study was conducted on a species (*Sinapis arvensis* L.) widespread in nature in Algeria particularly in the province of Tlemcen. This plant, even if it is undesirable in crop fields, can be useful in maintaining the biodiversity of the plant world. The goal set in this study was to treat the micro-propagation and germination of the species *Sinapsis arvensis* L. in synthetic environments, since these phenomena of growth and development in a sterile environment are poorly controlled. Our work deals successively with the following results: - The in vitro germination of *Sinapis arvensis* in different synthetic media, seeds taken of pods are harvested in stations in the Tlemcen region (Zenata, Beni-Ghanam and Rachgoun stations). They germinate differently; the germination rates vary with temperature and media used (Nutrient Agar and Potatoes Dextrose Agar, distilled water with NaCl at different concentrations, 1 g/L, 3 g/L and 5 g/L). This phenological phase appears to respond positively to the experimentation multiple conditions. - The percentage of germination was 75%. - Contamination by pathogens reaches 25%, despite the taken precautions (sterility of plant material, cleaning glassware, etc.).

Keywords: *Sinapis arvensis* L., Nutrient Agar, Potatoes Dextrose Agar Medium, Germination, Region of Tlemcen, Algeria

1. Introduction

Very effective in vitro culture remains a tool for biological and physiological research (Haicour, 2002).

Numerous studies have dealt with in vitro culture describing all used methods both in the animal and plant kingdoms (Harper and Benton, 1966; Dubey and Mall, 1972; Ungar, 1978; Koller and Hadas, 1982; Van Der Toorn and Ten Hove, 1982; Augéand al., 1989; Margara, 1989; Benchenafi and al., 2013). This is generally a method of growing plants in aseptic conditions (without fungi and bacteria) using quite complex culture media (hormones, sugars, vitamins, amino acids, mineral salts) that can be liquid, agar, or even solids with the use of vermiculite (Jay Allmand and Capelli, 1997).

We used the variability that occurs in vitro for an efficient plant according to some defined criteria; many other useful plant substances could be produced by tissue culture (Haines, 1995).

In vitro culture helps cultivate tissues or organs fragments isolated from a plant that can regenerate shoots, but also roots. This technique also provides isolated cells or even to regenerate a whole plant. In vitro culture produces the regenerative potential of a plant, to the cell totipotency which can result following this simple formula: 1 cell/unit = 1 whole plant (Jay-Allmand and Capelli, 1997).

Through this study we will perform an experiment (germination) with the species *Sinapis arvensis*, which is a Brassicaceae, and, according to Patrice (in Abdeldjelil, 2014), is considered as one of the most economically important ten plant families.

Particular attention will then be paid to the species vegetative stage, in this instance germination. In this work, we propose to vary the seeds sampling from the representative stations. Would the seeds meet these artificial food environments? Can these environments provide the best conditions for seed germination especially those belonging to *Sinapis arvensis* L.?

2. Methodology and Study Sites

2.1. Methodology

2.1.1. Disinfection of Equipment and Samples

In vitro culture technique requires great care in
maintaining cultures in asepsis conditions. Infections are either bacterial or fungal but these are most prevalent in the early cultures.

Disinfection of plant material is always difficult and unpredictable; the degree of infection of surface tissue is highly variable.

The disinfection method we used is the most common, which is to briefly immerse the sample in different solutions:
- Wash in running water,
- Immersion in a bleach solution,
- Three successive rinses with distilled water.

The seeds are disinfected according to the following protocol:
- Wash in running water,
- Soaking grains in ethyl alcohol at 70% for 20 to 25 seconds
- A bleach bath solution at 15 minutes,
- Three washes with distilled water at 10 minutes each.

The beads are then placed in petri dish; they are transplanted in the culture medium with sterile pliers.

The boxes are closed to prevent contamination; all manipulations are carried out under sterile hood.

A part of the dishes was stored at laboratory temperature (25°C) and another at 30°C and the last at 4°C for which we aim to test the effect of temperature on the germination.

2.1.2. Media Composition

The seeds are germinated in petri dishes containing 25 mL down different nutrient media at a rate of 10 seeds per box lined with blotting paper to produce seedlings for a period of 4 weeks.

To investigate this we used conventional medium containing sodium chloride (NaCl) at different concentrations (1 g/L, 3 g/L and 5 g/L) and two artificial media that are available in our laboratories (nutrient agar, and Potatoes Dextrose Agar) we experienced at temperatures of 4°C, 25°C and 30°C.

2.2. Study Sites

2.2.1. Zenata station

This station is located under the bridge in the national road RN 98 a few kilometers from the town of Zenata. It is located 1°27'West and 35°01' north, and the approximate altitude of the station is at 200 m.

2.2.2. Beni-Ghanam station

Located 1°17' West and 35°10' north and altitudinal level at 180 m, this second station is close to the national road 22.

2.2.3. Rachgoun station

This last station located west of Beni Saf and east of the Traras Mountains and is located at the mouth of Tafna Wadi near Rachgoun beach.

It is of 1°28' west and 35°17' north and its altitude is 54 m.

3. Results and Interpretations

In vitro culture technique requires great care in maintaining cultures aseptically.

When we have infected cultures, this may have various causes; it may be a fungus (mold) or a bacterium. If it is a fungus, we can see a mycelial development which has a whitish, greenish or grayish texture. If it is a bacterium, then
one can see a fog/veil of milky appearance, developed inside the medium and at the surface. If the infection starts from the contact area between the tissue and the medium, then the tissues are the sources of the infection which can be either to air or to an inadequate sterilization of the medium, or a contamination of ambient air through the water condensation of the lid (Augé and al., 1989).

According to Boccon-Gibod (1984) quoted by Heller (1990), it is the explants that are the source of infection. There also may be infections because of improper handling or use of non-sterile equipment. Aseptic conditions are difficult to create/reach in a laboratory (Loukidi, 1998).

The percentage of infection found on hypocotyls appears to be due to incomplete sterilization of seed, probably because of deep fungal infection. There are some simple and reactive techniques that can be used to ensure aseptic working conditions (Herbert and al., 1993; Loukidi, 1998).

Germination in room conditions (25°C) and agar.

- Zenata station: during the first week there was no germination, beginning the second week only germination shows successive increases, and moved sharply to 70% in the fourth week.
- Beni-Ghanam station: germination snaps from the 2nd week, and then it increases steadily until the fourth week where it reaches 50%.
- Rachgoun station: during the first week no germination was recorded; it was only from the second to the 4th week that germination shows successive increases to reach 100%.

### Table 1. Number of germinated seeds in Nutrient Agar medium, (Room temperature: 25°C).

| Weeks Stations | 1st Week Number | 1st Week % | 2nd Week Number | 2nd Week % | 3rd Week Number | 3rd Week % | 4th Week Number | 4th Week % |
|----------------|----------------|------------|----------------|------------|----------------|------------|----------------|------------|
| Zenata         | 0              | 0          | 4              | 40         | 5              | 50         | 7              | 70         |
| Beni-Ghanam    | 0              | 0          | 2              | 20         | 3              | 30         | 5              | 50         |
| Rachgoun       | 1              | 10         | 2              | 20         | 10             | 100        | 10             | 100        |

### Figures 1. Number of germinated seeds in Nutrient Agar medium, (Room temperature: 25°C).

Germination in room conditions (30°C) and agar.

### Table 2. Number of germinated seeds in Nutrient Agar medium, with an average temperature at 30°C.

| Weeks Stations | 1st Week Number | 1st Week % | 2nd Week Number | 2nd Week % | 3rd Week Number | 3rd Week % | 4th Week Number | 4th Week % |
|----------------|----------------|------------|----------------|------------|----------------|------------|----------------|------------|
| Zenata         | 6              | 60         | 6              | 60         | 6              | 60         | 6              | 60         |
| Beni-Ghanam    | 0              | 0          | 0              | 0          | 0              | 0          | 0              | 0          |
| Rachgoun       | 0              | 0          | 0              | 0          | 0              | 0          | 0              | 0          |

### Figures 2. Number of germinated seeds in Nutrient Agar medium, with an average temperature at 30°C.
Germination at average temperature (30°C) and agar. This germination case is very surprising; only the seeds of Zenata station responded favorably with 60% (steady from the beginning to the end of the experiment).

**Table 3. Number of germinated seeds in Nutrient Agar medium. (Cold Temperature: 4°C).**

| Weeks | Stations   | Number | %   | Number | %   | Number | %   | Number | %   |
|-------|------------|--------|-----|--------|-----|--------|-----|--------|-----|
|       | Zenata     | 0      | 0   | 0      | 0   | 0      | 0   | 0      | 0   |
|       | Béni-Ghanam| 0      | 0   | 0      | 1   | 1      | 10  | 10     | 10  |
|       | Rachgoun   | 3      | 30  | 5      | 50  | 5      | 50  | 6      | 60  |

**Figures 3. Number of germinated seeds in Nutrient Agar medium. (Cold Temperature: 4°C).**

Germination in cold temperature (4°C) and agar. Like the latter study case, this germination is also very surprising as only Zenata station seeds responded favorably with 60% (steady from the beginning to the end of the experiment).

**Table 4. Number of germinated seeds in the PDA medium. (at room temperature, 25°C).**

| Weeks | Stations   | 1st Week | % | 2nd Week | % | 3rd Week | % | 4th Week | % |
|-------|------------|----------|---|----------|---|----------|---|----------|---|
|       | Zenata     | 7        | 70| 7        | 70| 10       | 100| 10       | 100|
|       | Béni-Ghanam| 3        | 30| 4        | 40| 4        | 40 | 4        | 40 |
|       | Rachgoun   | 5        | 50| 6        | 60| 8        | 80 | 8        | 80 |

**Figures 4. Number of germinated seeds in the PDA medium. (at room temperature, 25°C).**

Germination in room conditions (25°C) and PDA medium.
- Zenata station: during the first and the second week we had 70% of germination, this rate reaches 100% in the last two weeks.
- Beni-Ghanam station: during the first week we have 30% of germination; from the 2nd to the 4th week it stabilizes at 40%.
- Rachgoun station: during the first week, germination starts at 50%, it is only in the second and 4th week that germination shows successive increases, we went from...
60% to 80%.

Table 5. Number of germinated seeds in the PDA environment. (Average temperature 30°C).

| Weeks | Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|-------|----------|----------|----------|----------|----------|
|       | Number   | %        | Number   | %        | Number   | %        |
| Zenata| 5        | 50       | 6        | 60       | 6        | 60       |
| Béni-Ghanam| 10       | 100      | 10       | 100      | 10       | 100      |
| Rachgoun| 4        | 40       | 5        | 50       | 5        | 50       |

Germination took place in an average temperature (30°C) and PDA medium.

- Zenata station: during the first week, germination rate is 50%, this percentage has stabilized at the 2nd week. During the third and fourth week we see significant increases (50% to 60%).
- Beni-Ghanam station: during the 4 weeks, we record a maximum germination which already appears at the first week (100%).
- Rachgoun station: after a sudden direct 40% rise, germination is stable up to week 4 and shows a 50% small increase.

Table 6. Number of germinated seeds in the PDA environment. (Cold temperature: 4°C)

| Weeks | Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|-------|----------|----------|----------|----------|----------|
|       | Number   | %        | Number   | %        | Number   | %        |
| Zenata| 0        | 0        | 1        | 10       | 2        | 20       |
| Béni-Ghanam| 0 | 0 | 0 | 0 | 1 | 10 |
| Rachgoun| 0        | 0        | 1        | 10       | 2        | 20       |

This germination is conducted at cold temperature (4°C) in PDA medium.

- Beni-Ghanam station: germination takes place only at the fourth week with a very small percentage (10%).
- Rachgoun station: germination begins in the 2nd week (10%), stabilizes during the third, ending ultimately the...
4th week with a percentage of 20%.

**Table 7.** Number of germinated seeds in a medium with salt concentration (1 g/L of NaCl) at room temperature (25°C).

| Weeks | Stations       | 1st Week |     | 2nd Week |     | 3rd Week |     | 4th Week |     |
|-------|----------------|----------|-----|----------|-----|----------|-----|----------|-----|
|       | Number | %         |     | Number | %     | Number | %     | Number | % |
| Zenata| 0      | 0         | 10  | 100     | 10   | 100      | 10   | 100      | 10  |
| Beïn-Ghanam | 0 | 0         | 0   | 0       | 0    | 0       | 0    | 0       | 0   |
| Rachgoun | 0 | 0         | 10  | 100     | 10   | 100      | 10   | 100      | 10  |

**Figures 7.** Number of germinated seeds in a medium with salt concentration (1 g/L of NaCl) at room temperature (25°C).

Seed germination in a medium with a salt concentration (NaCl) 1 g/L in room conditions (25°C) evolves in stations as follows:
- Zenata station: germination starts from the second week by showing a maximum (100%).
- Beni-Ghanam station: no sprouting here from the beginning to the end of the experiment.
- Rachgoun station: germination followed the same pattern as that of the Zenata station.

**Table 8.** Number of germinated seeds in a medium with salt concentration (1 g/L of NaCl) at medium temperature (30°C).

| Weeks | Stations       | 1st Week |     | 2nd Week |     | 3rd Week |     | 4th Week |     |
|-------|----------------|----------|-----|----------|-----|----------|-----|----------|-----|
|       | Number | %         |     | Number | %     | Number | %     | Number | % |
| Zenata| 0      | 0         | 0   | 0       | 0    | 0       | 0    | 0       | 0   |
| Beïn-Ghanam | 0 | 0         | 0   | 0       | 0    | 0       | 0    | 2       | 20  |
| Rachgoun | 0 | 0         | 1   | 10      | 1    | 10      | 2    | 20      |     |

**Figures 8.** Number of germinated seeds in a medium with salt concentration (1 g/L of NaCl) at medium temperature (30°C).

Seed germination in a medium with a 1 g/L of NaCl concentration and at an average temperature of 30°C:
- Zenata station: no sprouting is obtained in the four weeks of experimentation.
- Beni-Ghanam station: we record germination only in the fourth week, and that remains however low (20%).
- Rachgoun station: germination begins in the second week, remains steady in the third to finish at a rate of 20%.
the last week.

**Table 9.** Number of germinated seeds in a medium with 1 g/L salt concentration (NaCl) in cold temperature (4°C).

| Weeks | Stations    | 1<sup>st</sup> Week | 2<sup>nd</sup> Week | 3<sup>rd</sup> Week | 4<sup>th</sup> Week |
|-------|-------------|-----------------------|-----------------------|-----------------------|-----------------------|
|       | Number      | %                     | Number                 | %                     | Number                 | %                     | Number                 | %                     |
|       | Zenata      | 0                     | 0                     | 0                     | 0                     | 1                     | 10                    | 2                     | 20                    |
|       | Béni-Ghanam | 0                     | 0                     | 0                     | 0                     | 3                     | 0                     | 3                     | 30                    |
|       | Rachgoun    | 0                     | 0                     | 6                     | 60                    | 7                     | 70                    | 8                     | 80                    |

**Figures 9.** Number of germinated seeds in a medium with 1 g/L salt concentration (NaCl) in cold temperature (4°C).

Seed germination in a medium with 1 g/L (NaCl concentration) in cold temperature (4°C) showing the following changes:

- Zenata station: germination starts the third week with 10%, and then shows a change in the fourth week with 20%.
- Béni-Ghanam station: germination is recorded until the fourth week with a comparatively low (30%).
- Rachgoun station: germination snaps during the second week (60%), and increases in the last two with 70 and 80% respectively.

**Table 10.** Number of germinated seeds in a medium with 3g/L salt concentration (NaCl) at room temperature (25°C).

| Weeks | Stations    | 1<sup>st</sup> Week | 2<sup>nd</sup> Week | 3<sup>rd</sup> Week | 4<sup>th</sup> Week |
|-------|-------------|-----------------------|-----------------------|-----------------------|-----------------------|
|       | Number      | %                     | Number                 | %                     | Number                 | %                     | Number                 | %                     |
|       | Zenata      | 0                     | 0                     | 0                     | 0                     | 0                     | 0                     | 0                     |
|       | Béni-Ghanam | 0                     | 0                     | 0                     | 0                     | 0                     | 0                     | 0                     |
|       | Rachgoun    | 0                     | 0                     | 10                    | 100                   | 10                    | 100                   | 10                    | 100                   |

**Figures 10.** Number of germinated seeds in a medium with 3g/L salt concentration (NaCl) at room temperature (25°C).

Seed germination in a medium with a 3g/L salt concentration (NaCl) in room conditions (25°C) allows to note:

- Only Rachgoun station shows a maximum germination from the second week (100%), and of course remains steady during the other weeks.
The other two stations (Beni-Ghanam and Zenata showed no germination.

Table 11. Number of germinated seeds in a medium with 3g/L salt concentration (NaCl) at 30°C temperature.

| Weeks Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|----------------|----------|----------|----------|----------|
|                | Number   | %        | Number   | %        |
| Zenata         | 0        | 0        | 20       | 20       |
| Beni-Ghanam    | 0        | 0        | 0        | 0        |
| Rachgoun       | 0        | 0        | 0        | 0        |

Figures 11. Number of germinated seeds in a medium with 3g/L salt concentration (NaCl) at 30°C temperature.

Seed germination in a medium with salt concentration (NaCl) 3g/L room conditions (30°C) allows the following remarks:
- Zenata station: germination snaps the 2nd week with 20% and this rate remains steady during the other weeks.
- Beni-Ghanam station: no germination seen at this station.
- Rachgoun station: germination takes place only from the third week (10%) and ends with 40% in the 4th week.

Table 12. Number of germinated seeds in a medium with 3 g/L salt concentration (NaCl) at cold temperature (4°C).

| Weeks Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|----------------|----------|----------|----------|----------|
|                | Number   | %        | Number   | %        |
| Zenata         | 0        | 0        | 20       | 70       |
| Beni-Ghanam    | 0        | 0        | 30       | 50       |
| Rachgoun       | 0        | 0        | 10       | 50       |

Figures 12. Number of germinated seeds in a medium with 3 g/L salt concentration (NaCl) at cold temperature (4°C).

Seed germination in a medium with 3g/L salt concentration (NaCl) in cold temperature (4°C) allows to note:
- Zenata station: germination starts the second week with 20%, rises to 70% in the third to finish with a maximum (100%) in the last week.
- Both stations (Beni-Ghanam and Rachgoun): germination begins the second week with respective
percentages of 10 and 30% and ends with an average of 50%.

Table 13. Number of germinated seeds in a medium with 5 g/L salt concentration (NaCl) at room temperature (25°C).

| Weeks Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|----------------|----------|----------|----------|----------|
| Zenata         | 0        | 0        | 0        | 0        |
| Béni-Ghanam    | 0        | 0        | 0        | 0        |
| Rachgoun       | 0        | 0        | 0        | 0        |

Unfortunately, seed germination in a 5 g/L salt concentration (NaCl) medium at room conditions (25°C) does not show any germination in the three stations.

Table 14. Number of germinated seeds in a 5 g/L salt concentration (NaCl) medium at average temperature (30°C).

| Weeks Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|----------------|----------|----------|----------|----------|
| Zenata         | 0        | 0        | 0        | 3        |
| Béni-Ghanam    | 0        | 0        | 0        | 0        |
| Rachgoun       | 0        | 0        | 4        | 0        |

Seed germination in a 5 g/L salt concentration (NaCl) medium at average temperature (30°C) shows that germination takes place at the end of the experiment with 30% in Zenata station and 40% in Rachgoun station.

Table 15. Number of germinated seeds in a medium with a 5 g/L salt concentration (NaCl) at cold temperature (4°C).

| Weeks Stations | 1st Week | 2nd Week | 3rd Week | 4th Week |
|----------------|----------|----------|----------|----------|
| Zenata         | 2        | 20       | 8        | 80       |
| Béni-Ghanam    | 2        | 20       | 3        | 30       |
| Rachgoun       | 10       | 100      | 10       | 100      |
Seed germination in a medium with a 5 g/L salt concentration (NaCl) at cold temperature (4°C) leads us to notice:

- Zenata station: during the 2nd week we had a sudden sprouting at 80% which remained steady at this level until the end of the experiment.
- Beni-Ghanam station: during the first week there was a germination at 20%, which rose and remained steady at 30% until the end of the experiment.
- Rachgoun station: we have an excellent seeds response, to the point that we recorded a total germination (100%), representing a spectacular start of the germination.

4. Conclusion

Regarding this vegetative stage; it is a phenological stage in the life of plants (Sinapis arvensis germination seeds). To study this phenomenon we used two artificial media (Nutrient Agar and Potatoes Dextrose Agar) as well as distilled water to which we added different concentrations of NaCl (1 g/L to 5 g/L).

From the results obtained, the temperature has a significantly effect on this phenomenon; germination remains possible while the cold (4°C) inhibits the germination in an agar medium.

However, one must note that the germination conditions procedure was respected; microbial proliferation has not been too expressed.

As recommendations in the laboratory work, germination tests should be performed in maintaining well-defined conditions, because the presence of a single bacterium or fungus can trigger an invasion of the culture medium. That is why it is better to respect the rules, even simplify them. It would be necessary to ensure that all sterilized instruments are deposited in the sterile area, so as to have the slightest thing to do during handling and in sterile air.

References

[1] Abdeljalil A., 2014 – Quelques aspects germinatifs, rhizogéniques et écologiques chez Sinapis arvensis L. dans la région de Tlemcen. Mem. Mast. Ecol. Univ. Tlemcen, 139 p.
[2] Augé R., Beauchesne G., Boccon-Gibod J., Decourtye L., Digat B., Jalouzot R., Minier R., Morand J., Reynoard J. et Struulu D., 1989 - La culture in vitro et ses applications horticoles. 3ème édition revue, corrigée et augmentée. Ed Tech. Doc. Lavoisier 225 p.
[3] Benchefi Lachachi S., Benabadji N., Bemansour D., 2013 – Contribution to the study of Lygeum spartum L. germinative hostties in the south region of Tlemcen (Western Algeria). Environment Reserch Journal 7(2) ISSN 1994-5396: 20-24.
[4] Boccon-Gibod J., 1984 - Régénération du Crosne du Japon (Stachys sieboldii Mig.) par culture de méristème : multiplication et conservation in vitro des clones. In : Congrès sur l’application de la culture in vitro à l’amélioration des plantes potagères. ECARPIA, section légumes, Versailles. 31-41.
[5] Dubey P.S and Mall L.P., 1972 – Ecology of germination of weed seeds. I. Role of temperature and depth of burial in soil. Oecologia, 10: 105-110.
[6] Haicour R., 2002 - Biotechnologie végétale : technique de laboratoire. Ed Tec et Doc. Montréal. AUF, 2002 (université francophones ISBN 2-7430-0560-2). 275p.
[7] Haines H., 1995 - The ecology of running waters. Liverpool University Press, Liverpool. 555p.
[8] Harper J.L and Benton R.A., 1966 – The behavior of seeds in soil. II. The germination of seeds on the surface of water supplying substrate. J. Ecol. 54: 151-166.
[9] Hebert Y., Lefort-Buson M. et Damaerval M., 1993 - Les outils d’évaluation de la diversité génétique. Agronomie. 9(3): 173-178.
[10] Heller, 1990 - Physiologie végétale et développement 4ème Éd. Mass. Et Cie, 270p.
[11] Jay Allmand C., Capelli P. and Cornu D., 1992 - Root development of in vitro hybrid walnut microcutting in vermiculite containing gelrite medium. Station d’amélioration des arbres forestiers. INRA, 45160. Ardon France. SciencIA horticultura. 51(3-4) : 335-342.
[12] Jay Allmand C. et Capelli P., 1997-La multiplication végétative in-vitro, base méthodologique. D. E. A. Ressources génétiques et Amélioration des plantes. INA Paris Grignon. 101p.
[13] Koller D and Hadas Y, 1982 – Water relations in the germination of seeds. In: Lange O.I., Osmond C.B and Ziegler H (Eds.) Encyclopedia of plant physiology. Springer-Verlag, Berlin. 12(B): 401-431.

[14] Loukidi N., 1998 - Contribution à une étude morpho-histométrique de *Malva sylvestris* L. dans la région de Tlemcen. Mém. DES Physio. Vég. Univ. Tlemcen, 120 p.

[15] Margara F., 1989 - Bases de multiplication végétative : les méristèmes et l’organogénèse. Ed INRA, Paris. 262 p.

[16] Ungar I. A., 1978– Halophyte seed germination. *Bot. Rev.*, 44:233-264.

[17] Van Der Toorn J. and Ten Hove H. J., 1982 – On the ecology of *Cotula coronopifolia* L. and *Ranunculus sceleratus* L. II. Experiments on germination, seed longevity, and seedling survival. Acta *œcologica. œcol.Plant.* Vol. 3 (17), n°4. 409-418.