Effects of space environment on embryonic growth up to hatching of salamander eggs fertilized and developed during orbital flights

Lydie Gualandris-Parisol, David Husson, Alain Bautz, Danielle Durand, Paulette Kan, Christian Aimar, Hervé Membre, Anne-Marie Duprat, Christian Dournon

1 Center of Developmental Biology UMR 5547 CNRS- Paul Sabatier University, 118 Route de Narbonne, 31062 Toulouse cedex, France.
2 Laboratory of Experimental Biology-Immunology, EA 3442: Genetic, Signaling, Differentiation, Henri Poincaré University -Nancy-1, B.P. 239, 54506 Vandoeuvre-les-Nancy cedex, France.

Abstract In vertebrates, only few experiments have been performed in microgravity to study the embryonic development from fertilization. To date, these concern only amphibian and fish. We report here a study on the embryonic development of Pleurodeles waltl (urodele amphibian) eggs oviposited in microgravity. The experiment was performed twice on board the Mir space station and the data obtained included video recording and morphological, histological and immunocytological analyses. The data confirm that the microgravity conditions have effects during the embryonic period, particularly during cleavage and neurulation, inducing irregular segmentation and abnormal closure of the neural tube. Moreover, we observed several abnormalities hither to undescribed corresponding to cortical cytoplasm movements, a decrease of cell adhesion and a loss of cells. These abnormalities were temporary and subsequently reversible. The young larvae that hatched during the flight displayed normal morphology and swimming behavior after landing. The results obtained in the urodele Pleurodeles waltl are in accordance with those observed earlier in the anuran Xenopus laevis and in the fish Oryzias latipes.

Key words: ChAT, cleavage, neurulation, urodele, weightlessness.

Introduction
In vertebrates, the only experiments to date on fertilization and embryonic development in microgravity (µG) have been performed with amphibian and fish. Although dependent on experimental conditions, early development was perturbed.

In anurans, whose fertilization is external and monospermatic, the authors relied on artificial fertilization. In 1968, during the 11-hour Gemini mission, apparently normal cleavage stage Rana pipiens embryos were obtained from eggs fertilized on the launching pad (Young & Tremor 1968). Working on Xenopus laevis and using automatic hardware, Ubbels et al. (1989) first obtained artificial fertilization and the beginning of development in microgravity, on board sounding rockets. Under the same conditions, and then on board American space shuttles, she described abnormal blastulae and gastrulae with a thick blastocoel roof (Ubbels et al. 1994; Ubbels 1997). Souza et al. (1995) obtained fertilization and development of Xenopus tadpoles on board a space shuttle. Oocytes were fertilized in vitro with spermatozoa previously prepared on the ground. Ovulation of oocytes was induced by hormone injection during the flight, and fertilized eggs were then put on plates in microgravity or in a centrifuge permanently rotating at 1G. In flight, embryos were fixed for histological study or kept alive for rearing. Two-cell µG-embryos, gastrulae, neurulae and swimming tadpoles all appeared externally normal. However, the young µG-gastrulae had a thicker blastocoel roof than those developing under 1G conditions (Black et al. 1996).

In urodeles, the so-called AstroNewt experiments that used Cynops pyrrhogaster were performed on board the American shuttle Columbia on the Second International Microgravity Laboratory (IML-2) in 1994 and on board a Japanese Space Flyer Unit put in orbit in 1995 and retrieved by an U.S.A. Space Shuttle in 1996. The aim of these experiments was to study the development of urodele eggs laid by preinseminated females automatically injected or hormonally treated to trigger ovulation in microgravity conditions. Some females died during the flights but others laid about 71 eggs that developed into embryos judged as normal by examination of video images taken in orbit (Mogami et al. 1996; Yamashita et al. 2001).


During the 1990-1999 decade, research into amphibian development in space was particularly active for two main reasons; first, amphibians are a popular model in embryology and developmental biology, second, they can be reared on board space vehicles more easily than...
mammals. Our experiments were performed with *Pleurodeles waltl*. In comparison with *Xenopus laevis*, the interest of *P. waltl* is that the embryonic and larval development rates are slow and thereby facilitating analysis of various developmental stages. For example, at 18°C, the hatching stage is reached after 2 and 12 days in *Xenopus* and *Pleurodeles*, respectively. The goal of our project was to answer three questions. Can fertilization occur normally on board a space station? If so, is subsequent embryonic development normal in microgravity? The further development and reproduction be normal or not after return to Earth?

We performed two duplicate experiments on board the Mir space station. Control experiments were synchronously performed in a laboratory on the ground using the same hardware as that on board Mir. The first experiment took place during the 1996 Cassiopeia mission from August 17 to September 2, and the second one during the 1998 Pegasus mission from January 29 to February 19. In our experiments named ‘FERTILE’, the spermatozoa were preserved in the cloacal pelvic glands of preinseminated *P. waltl* females. During egg laying, they crossed the jelly coat and penetrated the oocytes in a natural manner. In a previous publication, we demonstrated that the development obtained on board Mir was the consequence authentic fertilization and not of parthenogenesis or gynogenesis (Aimar *et al.* 2000). We also showed that the subsequent larval development, metamorphosis and reproduction of animals oviposited and developed up to hatching stages in microgravity occurred normally (Dournon *et al.* 2001).

The aim of the present paper is to present statistically significant data concerning the embryonic development that occurred after fertilization obtained during space flights, up to the hatching stage. While confirming previous results, we observed abnormalities that had not been described earlier.

**Material and Methods**

**Animals:** All the *Pleurodeles waltl* (urodele amphibian) derived from the standard strain of the laboratory. The Cassiopeia mission took place outside the natural reproductive season of this salamander. Consequently, to keep alive the greatest possible number of spermatozoa in the cloacal glands after a natural mating, the inseminated females were maintained at 8°C for 12-16 weeks before the launch. The Pegasus mission occurred during the reproductive period. The inseminated females were maintained until the launch at room temperature (13-16°C) for 2-8 weeks after a natural mating. Embryos and larvae were staged according to the table of development of Gallien and Durocher (1957). The animals were treated according to the principles expressed in the Declaration of Helsinki and the Council Directive of the European Communities on the Protection of Animals Used for Experimental and Other Scientific Purposes (*L358/1, November 24, 1980*).

**The Fertile instrument:** An instrument named ‘Fertile’ was developed in collaboration with the French space agency (CNES) and the French manufacturer COMAT (Husson *et al.* 2001). It was an incubator set at 18 ± 1°C, connected to an air circulation system and containing a) a video camera to film the development of µG-embryos, b) three egg-laying boxes and two trays for the storage of egg-boxes placed in microgravity. Some egg-boxes included a capsule of fixative liquid, others were used to keep embryos alive. A ‘delrim’ grid separated the 14.85 ml water and egg compartment and the 17.93 ml air compartment. c) a centrifuge running at 1G on board Mir for the storage of egg-boxes at 18 ± 1°C. The arm of the centrifuge is 116 mm long (from the axis to the mid-central point in the egg box). d) nine radio-dosimeters.

**Measurement of radiation:** During the 17-day Cassiopeia space flight mission, the cumulated dose of cosmic and solar radiations was 350 ± 6 mrad and 10 ± 1 mrad on board Mir and on earth, respectively. During the 22-day Pegasus space flight mission, the dose was 457 ± 7 mrad and 15 ± 1 mrad on board Mir and on ground, respectively. The eggs and the embryos were less exposed because they were fertilized and developed after the beginning of the flight. On board Mir, the first embryos had the remaining 12 of 16 and 13 of 21 days, respectively, of development in microgravity and exposure to space radiations.

**General protocol:** For each launch, a thermally controlled container (CTA, in French: ‘Container de Transport Aller’) was used for the transportation of six *P. waltl* females in three boxes toward the Mir station. The CTA was set at 18 ± 2°C, and an air pomp ventilated (0.2 liter/min) the three boxes of transport. In each box, two adults were maintained without free water, but a damp towel covered the two largest internal surfaces and moisturized their skin. The trip toward the Mir station was two days long. During this period, the crew accessed to the animals. In flight, the protocol consisted in obtaining laying of fertilized eggs in two Fertile instruments, one on board Mir and the other in the ground laboratory. Egg laying was induced by an injection of 1.5 µl Luteinising Hormone-Releasing Hormone (LH-RH; Sigma, St Louis, MO, USA; 10 µg/ml in physiologic medium) into females that have been previously inseminated on the ground (Aimar *et al.* 2000). All the eggs were oviposited in microgravity, collected just after laying and distributed in twin boxes (about 20 eggs per box) with scissors and forceps. One box was stored on trays in ambient gravity and the other was placed in the centrifuge. The eggs were randomly placed in the centrifuge 1-4 hrs after laying. Most eggs of the twin batches were fixed at different times of the embryonic development. The others were kept alive. Due to the absence of a freezer on board Mir and of electricity for the experiments during the return of the Soyuz vehicles to Earth, we could not perform molecular analysis of deep-frozen embryos. For the return trip, the biological materials
were on board the Soyuz at the most one day. Up to
dedocking, the temperature increased progressively from
18 to 22 °C. From dedocking up to landing, the trip was
about six hours long and the temperature increased from
22 to 25 °C. After landing, the living animals were put at
18 °C. A similar profile was synchronously applied but the
control experiments performed in ground laboratory were
two days shifted. Effects of vibrations and shocks have
been previously tested on animals and materials (Husson
et al. 2001).

Fixing of the embryos; For the two space missions on
board MIR, embryos in their jelly coat developed in
hermetic egg-boxes at 18 °C in 8 ml of sterile Steinberg
rearing medium (Dournon et al. 2001). The formaldehyde
fixative was contained in capsules located in the egg-boxes.
A manual action distributed 500 µl of a 37 % formaldehyde
solution to give a final concentration of 2.2 %. Embryos
were kept in the fixative up to the end of the flights, and
then transferred to a 3.5 % formaldehyde solution. The same
protocol was applied to the ground control embryos.

Histological analyses and immunolocalizations; After
landing, all the fixed embryos were photographed. For
histological examinations, some were dehydrated in
ethanol, embedded in paraffin, cut into 8 µm sections, and
stained with safranin methyl blue orange G or hemalum
eosin. To perform immunolocalizations, other embryos
were incubated in increasing concentrations of sucrose
solutions (5, 10, 15, 20, 25 and 30 %), embedded in
Cryomount (Micron), frozen in isopentane in liquid
nitrogen and cut into 14 µm sections. Different primary
antibodies were used to compare the differentiation of
organs in µG- and 1G-embryos: anti-GFAP, a polyclonal
antibody directed against a glial fibrillary acidic protein
(Dakoppats); anti-NC1, a monoclonal antibody directed
against a neuronal cell membrane glycoprotein and used
as a specific neuronal cell marker in Pleurodeles, (Saint-
Jeannet et al. 1990); anti-MF20, a monoclonal antibody
directed against a sarcomere myosin of striated muscle
(Hybridoma bank); anti-GABA, a polyclonal antibody
directed against gamma aminobutyric acid-containing
neurons (Pituello et al. 1989). Two fluorescein (FITC)-
or rhodamine (TRITC)-conjugated secondary antibodies were
applied using standard protocols. The immunolocalizations
were performed with confocal (LSM 410, Zeiss) or
epifluorescence (Leitz Dialux 20) microscopy.

Choline acetyltransferase (ChAT) assays; Live hatching
larvae were cooled 8-10 hrs after the landing of each space
mission. They were 8.5 and 10.5-days old, respectively.
ChAT activity was measured in the presence of tritiated
acetyl Coenzyme A (5.8 mCi/mol, Amersham) using a
previously described method (Duprat et al. 1985). The
results were expressed as pmol/min/embryo.

Statistical analysis; To compare both values and
percentages of embryos, we used the χ² test with the Yates
correction (χ²c). The limit of significance for all the tests
was 3.841 for 1 degree of freedom and α = 0.05. To compare
the thickness of the blastocoel roof, Student’s t test was
used. To compare the ChAT activity, we used the Mann-
Whitney test.

Results

1 - Embryos developed on board MIR and on the ground

On board MIR, 10 out of 12 inseminated females, reared
table 1 embryos layed on board the Mir station or in the ground laboratory. At left, number (No) and percentage (%) of developed
embryos fixed in flight at different stages of development. At right, number (No) and percentage (%) of embryos obtained alive after
delivery of the samples on ground. Some embryos were used for histological analyses and ChAT assays

<table>
<thead>
<tr>
<th>Embryos fixed during the flight</th>
<th>Embryos kept alive during the flight</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of eggs used to study the embryonic development</td>
<td>No of developed and fixed embryos</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>-----------------------------------</td>
</tr>
<tr>
<td>Cassiopeia mission</td>
<td></td>
</tr>
<tr>
<td>In µG on board MIR</td>
<td>504</td>
</tr>
<tr>
<td>In 1G MIR centrifuge</td>
<td>147</td>
</tr>
<tr>
<td>In 1G on ground</td>
<td>527</td>
</tr>
<tr>
<td>Pegasus mission</td>
<td></td>
</tr>
<tr>
<td>In µG on board MIR</td>
<td>719</td>
</tr>
<tr>
<td>In 1G MIR centrifuge</td>
<td>173</td>
</tr>
<tr>
<td>In 1G on ground</td>
<td>680</td>
</tr>
<tr>
<td>Total missions</td>
<td></td>
</tr>
<tr>
<td>In µG on board MIR</td>
<td>1223</td>
</tr>
<tr>
<td>In 1G MIR centrifuge</td>
<td>320</td>
</tr>
<tr>
<td>In 1G on ground</td>
<td>1207</td>
</tr>
</tbody>
</table>
at 18°C, provided embryos, which developed in μG and in 1G. Of 12 ground control females, 11 provided embryos that developed in 1G. The percentage of development was defined as the number of developed embryos at the 2-cell stage or later stages divided by the total number of eggs. The numbers and percentages of development of in-flight and on-ground embryos were similar for each space mission. The Cassiopeia mission took place outside the reproductive period, whereas the Pegasus mission occurred during the reproductive period and the percentage of development was therefore optimal (Table 1). The timing of development was established by observation of the fixed embryos, annotations of the crew at each fixation time and video recording.

2 - Cleavage period
This period includes stages 1 to 7. Embryos at all these stages were obtained in flight.

Effects of microgravity on the inner cytoplasm of eggs. Sections of μG-embryos at the beginning of cleavage were examined under light microscopy. In these eggs, yolk platelets were distributed along an animal pole-vegetal pole axis in a gradient of increasing size, the smallest ones laying at the animal pole, as in 1G-egg controls. No striking modification of the vitelline gradient was observed in the different replicates. In addition, the cortical cytoplasm underneath the cell membrane of μG-eggs was yolk free and had the same constant thickness as that of 1G-control eggs.

Cleavage. At the beginning of cleavage, three kinds of abnormalities were observed in μG-embryos. A given embryo expressed one or several abnormalities and all possible combinations were observed (Fig. 1A to D and L to O; Table 2). The aspect and position of the pigmented animal area were modified and were unrelated to the orientation of the cleavage planes. This modification was observed in 34.4%, 13.0% and 11.9% of μG-, 1G-on board and 1G-ground embryos, respectively. The difference between μG- and 1G-centrifuge embryos was not statistically significant ($\chi^2 = 3.694$) but was significant between μG- and 1G-ground embryos ($\chi^2 = 6.687$). There was no difference between 1G-centrifuge and 1G-ground embryos ($\chi^2 = 0.005$) (Table 2).

The cleavage was irregular. For example, three, five or seven cells were observed in the animal hemisphere. This was seen in 31.6%, 7.7% and 2.3% of μG-, 1G-on board and 1G-ground embryos, respectively. The difference between μG- and 1G-centrifuge embryos was not statistically significant ($\chi^2 = ...$)
Gualandris-Parisot, L. et al.

1.777) but significant between µG- and 1G-ground embryos (χ² 10.728). There was no difference between 1G-centrifuge and 1G-ground embryos (χ² 0.003).

The intercellular space was enlarged at the animal surface of the embryo, indicating a decrease of cell adhesion. This was observed in 40.4%, 10.7% and 1.5% of µG-, 1G-on board and 1G-ground embryos, respectively (Figs. 1D and 2A). The differences were statistically significant between µG- and 1G-centrifuge (χ² 6.101) or 1G-ground embryos (χ² 25.706). There was no difference between 1G-centrifuge and 1G-ground embryos (χ² 2.104).

At blastula stages, vitellus pellets were observed between the blastomeres of some µG-embryos, suggesting a rupture of the membrane of inner cells of the vegetal hemisphere (Fig. 1E). Small cells of the periphery were detaching (Fig. 1F and G). The 1G-controls had no such an abnormality. Histological observations of embryos at stage 7 indicated that the blastocoel was located inside the animal hemisphere, as it was in control embryos. The roof of the blastocoel, made up of two layers of cells, had an identical thickness: 194.0 ± 56.7 and 196.7 ± 82.7 µm for the µG- and 1G-blastulae (Student’s t test not significant p = 0.90).

3 - Gastrulation period

This period encompasses stages 8 to 12. Only embryos at stages 8 and 9 were obtained in flight. Histological observations showed that the blastocoel roof was thicker in µG- than in 1G-gastrulae at stage 8, 202.1 ± 36.2 and 130.0 ± 38.5 µm, respectively (Student’s t tests significant p = 0.0008). However, as in the case of blastula, the roof was always composed of two layers of weakly cohesive cells in µG-embryos (not illustrated).

4 - Organogenesis

Neurulation period

This period encompasses stages 13 to 21. Embryos at stages 16 to 21 were obtained in microgravity. All the µG-embryos without exception displayed normal bilateral symmetry. At stage 21, 81.3% of the µG-embryos showed an incomplete closure of the neural tube at the cephalic and trunk levels. During neural tube closure, cells were

<table>
<thead>
<tr>
<th>Period</th>
<th>µG-embryos on board MIR</th>
<th>1G-embryos centrifuged on board MIR</th>
<th>1G-ground control embryos</th>
<th>χ² test 1, 2, 3</th>
<th>Student t-test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cleavage Abnormalities stages 1 to 5</td>
<td>No of embryos</td>
<td>121</td>
<td>75</td>
<td>101</td>
<td></td>
</tr>
<tr>
<td>Abnormal pigmentation</td>
<td>No of cases</td>
<td>10/29</td>
<td>6/46</td>
<td>12/101</td>
<td>ns, ns</td>
</tr>
<tr>
<td>%</td>
<td>34.4 %</td>
<td>13.0 %</td>
<td>11.9 %</td>
<td>ns, ns, ns</td>
<td></td>
</tr>
<tr>
<td>Irregular cleavage</td>
<td>No of cases</td>
<td>12/38</td>
<td>1/13</td>
<td>1/43</td>
<td>ns, ns</td>
</tr>
<tr>
<td>%</td>
<td>31.6 %</td>
<td>7.7 %</td>
<td>2.3 %</td>
<td>ns, ns, ns</td>
<td></td>
</tr>
<tr>
<td>Enlarged intercellular space</td>
<td>No of cases</td>
<td>19/47</td>
<td>3/28</td>
<td>1/65</td>
<td>ns, ns</td>
</tr>
<tr>
<td>%</td>
<td>40.4 %</td>
<td>10.7 %</td>
<td>1.5 %</td>
<td>s, s, ns</td>
<td></td>
</tr>
<tr>
<td>Normal embryos stage 7</td>
<td>Blastocoel roof thickness</td>
<td>No of cases</td>
<td>16</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>µ m</td>
<td>194.0 ± 56.7</td>
<td>196.7 ± 82.7</td>
<td>ns</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gastrulation Abnormalities stage 8</td>
<td>No of embryos</td>
<td>48</td>
<td>19</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>Blastocoel roof thickness</td>
<td>No of cases</td>
<td>28</td>
<td>8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>µ m</td>
<td>202.1 ± 36.2</td>
<td>130.0 ± 38.5</td>
<td>s</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neurulation Abnormalities stages 16 to 21</td>
<td>No of embryos</td>
<td>37</td>
<td>21</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td>Incomplete neural tube closure</td>
<td>No of cases</td>
<td>13/16</td>
<td>1/12</td>
<td>0/10</td>
<td>ns, ns</td>
</tr>
<tr>
<td>%</td>
<td>81 %</td>
<td>8 %</td>
<td>0 %</td>
<td>s, s, ns</td>
<td></td>
</tr>
<tr>
<td>Tail bud Abnormalities stages 22 to 36</td>
<td>No of embryos</td>
<td>63</td>
<td>54</td>
<td>32</td>
<td></td>
</tr>
<tr>
<td>Small head</td>
<td>No of cases</td>
<td>16/63</td>
<td>10/54</td>
<td>3/32</td>
<td></td>
</tr>
<tr>
<td>%</td>
<td>25.4 %</td>
<td>18.5 %</td>
<td>9.4 %</td>
<td>ns, ns, ns</td>
<td></td>
</tr>
<tr>
<td>Ascite Stages 28 to 33 only</td>
<td>No of cases</td>
<td>6/30</td>
<td>4/19</td>
<td>1/16</td>
<td></td>
</tr>
<tr>
<td>%</td>
<td>20.0 %</td>
<td>21.1 %</td>
<td>6.2 %</td>
<td>ns, ns, ns</td>
<td></td>
</tr>
<tr>
<td>Total number of analyzed embryos</td>
<td>269</td>
<td>169</td>
<td>183</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* χ² test: 1, difference between µG- and 1G-embryos centrifuged on board MIR; 2, difference between µG- and 1G-embryos on ground; 3, difference between 1G-embryos centrifuged on board MIR and 1G-ground control embryos (ns, no significant; s, significant). ' Sampling of embryos developed in egg boxes from one or both missions.
Salamander development in weightlessness

shed from the neurctoderm and were observed outside the tube (Fig. 1H, I and P). In contrast, the cells of the epidermis that differentiated from ectoderm showed normal adhesion. In 1G-embryos, only 8.3 % of the 1G-centrifuge and 0 % of the 1G-ground embryos exhibited an abnormal neurulation (Table 2). The differences were statistically significant between μG- and 1G-centrifuge (χ² = 11.812) or 1G-ground embryos (χ² = 13.162). There was no difference between 1G-centrifuge and 1G-ground embryos (χ² = 0.007).

In the course of development, the epidermal cells acquired cilia, the movement of which progressively induced a rotation of the embryos within the egg membranes. Under natural conditions, this movement was very slow and could only be observed by time-lapse video-recordings. Under microgravity conditions, video-movies confirmed that rotation began at the embryonic stage 16 and occurred during the tail bud stages, just as it did in control embryos.

Tail bud stages up to hatching stages

This period encompasses stages 22 to 34/36.

Morphological aspects. Under microgravity conditions, the closure of the neural tube was completed at stage 31 but was abnormal at the cephalic level. Moreover, 25.4 %, 18.5 % and 9.4 % respectively of the μG- and 1G-centrifuged and 1G-ground embryos had a small head. The differences were not statistically significant between μG- or 1G-centrifuge (χ² = 0.450) or 1G-ground embryos (χ² = 2.475) and also between 1G-centrifuge and 1G-ground embryos (χ² = 0.674). As the μG-, 1G-centrifuge and 1G-ground embryos were fixed during the experiments, their subsequent development could not be followed. However, of the 16 fertilized eggs put in the video-recording box, 14 (87.5 %) reached the hatching stage just before the end of the space flight. Two neurulae were abnormal and became abortive tail bud embryos.

The development of cardiac ascites in Pleurodeles embryos was used as an indicator of poor rearing conditions. There were no statistically significant differences between μG-, 1G-centrifuge and 1G-ground embryos (χ² = 0.075; 2.70; 0.579) in the frequency of the appearance of ascites. At stages 24 to 32 (Fig. 1J, K), histological analysis indicated no differences between μG-embryos and 1G-centrifuged embryos in the central nervous system (CNS), eyes and ears, or in the morphology of the
head and trunk (Fig. 2B, C). Five hours after landing, six living μG-hatched-embryos were fixed. The central nervous system appeared normal. The developing brain subdivisions were structured, and the white matter was located at the periphery of the CNS. Moreover, the differentiation of spinal cord, pronephros and gut were also normal in comparison with 1G-embryos.

At stage 30, the video-recordings showed that the first spontaneous contractions of the trunk musculature occurred in μG- and 1G-ground embryos. At stage 32, after 180 ± 3 hrs of development, the first μG-embryos hatched even though the 1G-embryos hatched after 230 ± 3 hrs, at stage 33b. At stages 33/34, 10 hours after landing, the young-μG larvae observed with binocular microscope and video-recording displayed swimming behavior identical to that of controls, with no looping or somersaulting. Except for the hatching event, the timing of the embryonic development was the same for μG- and 1G-young larvae.

**Immunolocalizations.** GABA is an inhibitory transmitter in the CNS. Between stages 26 and 31, no cells stained for GABA were observed in μG- or in 1G-ground embryos, whereas at stages 33 and 34 marked cells were observed both in μG- and in 1G-embryos. In 1G-ground embryos, GABA was first detected at stage 32b-33. The labelled cells were located in the periphery of the spinal cord, in white matter and in cellular aggregates (6-10 cells) in the neural tube and in retinal neurons. GFAP immunolocalization was performed on 6 embryos at stages 28 to 34. In the spinal cord and at the cephalic level, the radial glia differentiated with an identical time-course in μG- and 1G-ground embryos. The radial extensions were located throughout the neural tube except in its dorsal and ventral walls. The glial buds were normally located and timed (Fig. 2D to F). Astrocytes visualized with anti-GFAP were distributed normally in the differentiating retina in all cases. Using anti-NC1 and phase-contrast microscopy, cell bodies and axons in the diencephalon and in the eyes were seen to be normally located and developed with identical kinetics in μG- and 1G-centrifuge embryos (Fig. 2B and C). Anti-MF20 revealed no delay or positional abnormality in the differentiation of the somites. At stage 32, organized striated myofibrils appeared with normal chronology in the ventral region of the somite (Fig. 2G).

**ChAT activity.** This enzyme is involved in the biosynthesis of acetylcholine, the neurotransmitter of cholinergic synapses in the CNS and PNS, and of neuromuscular junctions. ChAT displayed the same activity at stages 32b⁷ and 33b in μG-embryos (N = 5 and 3) and in 1G-ground controls (N = 9 and 8). In control embryos, ChAT activity was first detectable at stage 31; it increased up to stage 34 and remained constant thereafter until stage 36 (N = 4-5 embryos per stage) (Fig. 3).

**Discussion**

The percentages of development of the embryos on board Mir and on earth were weak for the Cassiopeia mission and normal in the case of the Pegasus mission (Table 1). Nevertheless, these were consistent with the preparation of females for the space flights and with the seasons at which the two space missions occurred. On the other hand, it must be pointed out that they was no significant difference in development between our two kinds of controls: the on board 1G-centrifuge and 1G-ground embryos (Table 2).

At the beginning of cleavage, microgravity had effects on embryos. In the animal hemisphere, the standard distribution of the cortical cytoplasm including pigment granules was disturbed for 34.4% of μG-embryos and cleavages were irregular for 31.6%. The data revealed a significant difference between μG- and 1G-ground embryos, but surprisingly, no significant difference between μG- and 1G-centrifuge Mir embryos. It must be pointed out that all on board embryos were obtained from eggs oviposited in microgravity, because the cosmonaut randomly placed the fertilized eggs in the centrifuge 1-4 hrs after laying. Consequently, microgravity was able to act on the eggs before the first cleavage (5-6 hrs) explaining, for both alterations in pigmented area and cleavage, the absence of statistical difference between μG- and 1G-centrifuge Mir embryos. In Pleurodeles, Aimar et al. (2000) demonstrated that gravitational forces modify the cell membrane of fertilized eggs. We hypothesize that such a phenomenon also occurs during subsequent development, thereby explaining the abnormalities observed in embryos oviposited in microgravity. Another hypothesis could be proposed. As no significant differences of abnormal pigmentation and irregular cleavage between μG- and 1G-embryos on board Mir, and significant differences were observed between μG-embryos on board Mir and 1G-embryos on earth, these results could suggest that some abnormalities observed were possibly caused by on board environments rather than by simple environmental factors.

In Pleurodeles, 31.6% of μG-embryos displayed irregular cleavage. In Xenopus, Souza et al. (1995) indicated that under microgravity conditions, the cleavage
furrow was positioned normally at the two-cell stage, but the mitotic asters showed a slight displacement toward the vegetal pole. Using a clinostat on the ground to simulate microgravity, Neff et al. (1993) and Yokota et al. (1994) observed that the third cleavage shifted toward the embryo equator. Consequently, microgravity has effects on embryos of both *Xenopus* and *Pleurodeles* at the beginning of cleavage.

In spite of some asymmetrical segmentations in *Pleurodeles*, all the μG-embryos subsequently acquired normal bilateral symmetry. Consequently, as for *Cynops pyrrhogaster*, the egg rotation that occurs on earth after fertilization is clearly not necessary in urodeles to determine the position of symmetrical plan (Yamashita et al. 2001). This result supports previous ones for fish in which gravity was shown not to be necessary for the dorsal/ventral axis orientation or subsequent development (Wacker et al. 1993; Ijiri 1995, 1997). It also agrees with previous studies in anurans (Black et al. 1996; Souza et al. 1995; Ubbels et al. 1989, 1994), in which sperm impact induces bilateral symmetry. However, this phenomenon cannot be involved in *Pleurodeles*, where fertilization is polyspermy.

During cleavage, the intercellular space was significantly enlarged in 40.4% of μG-embryos. This morphological aspect, which could be interpreted by a decrease in cell adhesion, was also observed at the beginning of gastrulation and during neurulation.

At the beginning of gastrulation in *Pleurodeles*, the blastocoel roof of 1G-embryos normally decreases in thickness. In contrast, the blastocoel roof of μG-embryos remained thick, like at late the blastula stage. The thickness of the μG-embryo roof was related to a reduction of adhesion between ectodermal cells and not to an increase of the number of cell layers, as observed by Ubbels et al. (1994) for μG-embryos of *Xenopus laevis* or with cell proliferation as proposed by Black et al. (1996). However, the data of Neff et al. (1993) who used a clinostat suggested that the thickening of the blastocoel roof of *Xenopus* embryos was due to a change in the latitude of the third cleavage plane rather than a modification of cell proliferation. Moreover, De Mazière et al. (1996) observed in μG-*Xenopus* that the blastocoel shifted toward the vegetal hemisphere. The difference between frogs and salamanders could be due to modifications of osmoregulation and volume of the blastocoel liquid (Ubbels et al. 1997).

During gastrulation in *Pleurodeles*, only young gastrulae were fixed in flight, and consequently, insufficient histological information was obtained on this important embryonic period. Two days after fertilization, organogenesis began with neurulation. Cell adhesion was also perturbed in μG-embryos and cells detached. The cellular morphogenetic movements of the forming neural tube occurred more slowly in μG-embryos than in 1G-controls and standard embryos. In contrast, epidermis differentiated normally in these embryos. A weakening of the neurectodermal cell junctions could cause a difference in cell adhesion, even if the ectoderm and epidermal cells acquired functional specific junctions, in time. At stage 16, resulting from a ciliary beating movement, the μG-embryos began to turn inside their fertilization membrane in the same way as in 1G-control embryos.

During the organogenesis that leads to hatching, although numerous tail-bud embryos appeared with a small head, the statistical differences between the percentages were not significant. Moreover, the percentage of development indicated that these embryos continued their development normally. This was confirmed by video-movies. Of the 16 developing μG-embryos, only two abnormal neurulae became abortive tail bud embryos. All the other embryos developed or regulated.

During organogenesis, the immunohistochemical analyses indicated no gross differences between μG- and 1G-embryos in the structural differentiation of organs such as the nervous system. These data are in accordance with previous results obtained during a 12-day space flight. Cultured *in vitro*, isolated neurectodermal and mesodermal cells differentiated in a normal manner into neurons, glial and muscle cells (Husson et al. 1997). Moreover, the spontaneous muscle contractions occurring before the hatching period and the swimming motor reflexes performed by the first mature neural circuit appeared at the same time in μG-embryos and 1G-controls.

On earth, the mean annual dose of received solar and cosmic radiations is about 150 mrad a year, and the admitted human 50% lethal dose is 450 rad. On the ground, the atmosphere protects life from most of the cosmic radiations, but on board a space station, the crew and the living material received all the radiations. During both space flights, the dose received was 30-35 time high than on the ground. The duration of development at 18°C, from egg laying up to the hatching, was comparable on board the space station and in the ground laboratory. During their further development up to adulthood and reproduction, no abnormality distinguished these animals from standard ones (Dournon et al. 2001). Consequently, it appears that the cosmic radiations did not perturb the development of the animals. This finding agrees with the absence of a significant difference between on board 1G-centrifuge and 1G-ground embryos (Table 2).

The present ontogenetic study in microgravity extends the results obtained with the anuran *Xenopus* (Ubbels et al. 1989, 1994; Souza et al. 1995; Black et al., 1996) and the fish *Medaka* (Ijiri 1995, 1997). In *Pleurodeles*, the observed abnormalities, cortical cytoplasm movements, decrease of cell adhesion and loss of cells, probably concern the cell membrane, as suggested by the results of Schatz et al. (1990), Claassen et al. (1996) and Aimar et al. (2000). Modifications or alterations of physico-chemical properties of the membrane could involve the cytoskeleton (Spooner et al. 1994; Tabony 1994; Lewis et al. 1998). It will be interesting to determine whether abnormalities involve modifications of cell membrane or cell junctions.

In conclusion, the data confirm that microgravity conditions have effects during the embryonic period and particularly during cleavage and neurulation of amphibians.
Although the early development was not strictly normal, as a consequence of embryological regulation phenomena, the young hatching larvae had a normal morphological aspect and swimming behavior after landing. Video-movies performed on board Mir clearly confirm this point.

Acknowledgments

We would like to thank Françoise Foulquier and Julian Smith from the Developmental Biology Center in Toulouse and Christiane Tankosic from the Experimental Biology-Immunology Laboratory in Nancy for technical assistance. We gratefully acknowledge all the CNES board and particularly Didier Chaput and Michel Viso, for engineering and management. We have the great pleasure to thank the French cosmonauts Dr. Claude André-Deshays and Lieutenant Colonel Léopold Eyhartz for efficient practical expertise. We are grateful to our Russian colleagues and particularly the RKK Energia ones for preparation and success of the space missions. This work was supported by grants of the French space agency, the Centre National d'Études Spatiales.

References


