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Examining the impact of multi-layer graphene using cellular and amphibian models

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Abstract
In the last few years, graphene has been defined as the revolutionary material showing an incredible expansion in industrial applications. Different graphene forms have been applied in several contexts, spreading from energy technologies and electronics to food and agriculture technologies. Graphene showed promises also in the biomedical field. Hopeful results have been already obtained in diagnostic, drug delivery, tissue regeneration and photothermal cancer ablation. In view of the enormous development of graphene-based technologies, a careful assessment of its impact on health and environment is demanded. It is evident how investigating the graphene toxicity is of fundamental importance in the context of medical purposes. On the other hand, the nanomaterial present in the environment, likely to be generated all along the industrial life-cycle, may have harmful effects on living organisms. In the present work, an important contribution on the impact of multi-layer graphene (MLG) on health and environment is given by using a multifaceted approach. For the first purpose, the effect of the material on two mammalian cell models was assessed. Key cytotoxicity parameters were considered such as cell viability and inflammatory response induction. This was combined with an evaluation of MLG toxicity towards Xenopus laevis, used as both in vivo and environmental model organism.

Introduction
Graphene is one-atom-thick planar sheet of sp²-bonded carbon atoms that are densely packed in a honeycomb crystal lattice [1]. Graphene can display different forms which are conventionally combined under the so-called graphene family materials (GFMs). GFMs comprise few-layer graphene (FLG), multi-layer graphene (MLG), graphene oxide (GO), reduced graphene oxide (rGO), graphene nanosheets, ultrathin graphite, graphene ribbons and graphene dots. FLG is constituted by 2–5 graphene layers, while GO and rGO are normally composed of a single layer [2]. Despite being the basic structural element of other carbon allotropes, including carbon nanotubes and fullerenes, graphene history is quite recent. In fact, the research group, guided by Geim and Novoselov, reported for the first time a method for the isolation of single-layer graphene from graphite only in 2004 [3]. From its discovery, graphene has been the object of great industrial interest. Such success is due to its exceptional physico-chemical properties such as
electronic, optical, thermal and mechanical which differ from the same material in bulk. Such properties can be tuned by choosing a particular type of graphene [4]. By taking advantage of its special characteristics, graphene-based nanotechnology represents nowadays an area of scientific research and industrial applications in full expansion [5, 6]. In fact, graphene was firstly exploited in material sciences and recently it showed to be a good candidate for the development of graphene-based electronics, photonics, composite materials, energy generation, energy storage and sensors [5–9]. Among all graphene forms, FLGs are easily produced in high yield [10–13]. Low-cost procedures have also been described for the fabrication of large area films of FLG for electronics and opto-electronic applications [14].

This rapid increase in production and applications of graphene implies its potential release into the environment, especially into the aquatic compartment, which usually concentrates all kinds of contaminants [15]. For example, graphene release could occur from the use of commercial products containing the material, from its degradation during use and from the waste disposal of such products. The possible ecosystem risks induced by graphene were described by Hu and Zhu [15]. What is the environmental impact that can be expected after graphene introduction into the aquatic environment? The material would first interact with abiotic compounds such as natural organic matter and other molecules naturally present in waters (inorganic and organic macromolecules, colloidal particles, etc) but also with living organisms. Being important components of the ecosystem, the responses of aquatic organisms to graphene are particularly critical for ecosafety [15]. However, the existing knowledge of the potential toxicity of GFMs towards aquatic organisms is still poor. A recent review presents what is known on the toxicity of GFMs towards the aquatic environment [16]. Most of these works deal with the toxicity of GO and rGO on different aquatic organisms (bacteria, crustaceans, nematodes and fishes). Only a few studies report toxicity evaluation of single- and MLG. The general conclusion is that the impact of graphene against aquatic organisms may not be as high as carbon nanomaterials such as fullerenes and carbon nanotubes [16].

On the other hand, the possibility to isolate graphene flakes by repeated mechanical exfoliation of graphite, provides a material with a high degree of purity, making it suitable also for biomedical applications [17]. In addition, its high surface area allows for the conjugation with bioactive molecules such as DNA, proteins, peptides or small drug molecules [17]. Single- and FLG, demonstrated their potential in drug/gene delivery, biosensing and imaging, antibacterial activity and tissue engineering [18–21]. It is evident how such applications are strictly dependent on graphene biocompatibility. In fact, its use for therapeutic purposes necessitates the demonstration of the absence of graphene-induced tissue damage or proinflammatory response. A large variety of studies can be found in literature reporting the toxicity findings on graphene in vitro and in vivo [22]. Despite this, the described results are sometimes in contrast. In fact, the biological response varies depending on graphene number of layers, lateral size, rigidity, hydrophobicity, dose administered and purity of the material. The use of diverse cellular models may also be responsible for paradoxical findings [22]. For this reason GFMs’ toxicological profile is little understood and more studies concerning specific graphene types are required to avoid unexpected harmful effects on human health.

The current work is aimed at better elucidating the impact of graphene mainly consisting of 2–20 layers (MLG) on health and environment. The boundary between MLGs and graphite nanoplatelets is not sharp, so we decided to identify our material as MLGs even if the number of layers exceeds 10 (see [2] for a proposed nomenclature describing the different forms of graphene). For this purpose, two different mammalian cellular lines (human epithelial cells or murine macrophages) were used and key acute toxicity parameters such as cell viability and induction of proinflammatory response upon exposure to MLG were examined. The importance of MLG purification from adsorbed solvents was also taken into account. The in vitro preliminary studies were combined to the in vivo evaluation of MLG toxicity on the aquatic amphibian model organism Xenopus laevis. We used normalized exposure conditions [23] to assess larval mortality, growth inhibition and genotoxicity. On the other hand, the amphibian larvae represent also a relevant environmental model of aquatic organism to study the ecotoxicity of nanoparticle [24–27].

Materials and methods

Materials

MLGs were generated through a simple, fast and low cost production protocol with high yield previously described [10]. The flakes are made of sheets with different lateral size. This specific structure is useful for several applications (i.e. in catalysis, battery, supercapacitors and others, where the stabilization of metal particles are important) and it is related to the preparation process [28]. MLGs obtained by this method consist mainly of 2–20 sheets that, occasionally, can go up to fifty (see supporting figure S1). The thicker flakes were subsequently separated from the sample by a sedimentation process in toluene. The high purity sample (MLG1) was thermally treated to eliminate potentially adsorbed toluene between the graphene layers (MLG2) [29]. The lateral size of the MLG sheets ranged from 1.2 to 5.4 μm, with the average centered at 2.3 μm (see supporting figure S2). MLG dispersions were prepared freshly prior to each cellular exposure by sonicating the material in cell
culture medium containing 10% of fetal bovine serum (FBS), at a concentration of 100 μg ml⁻¹. The samples were diluted to the proper concentration before use.

Transmission electron microscopy
The morphology and number of MLG flakes were examined by high-resolution transmission electron microscopy on a Topcon 002B-UHR microscope working with an accelerated voltage of 200 kV and a point-to-point resolution of 0.17 nm. Prior to the analysis, the sample was ultrasonically dispersed in ethanol during 5 min and a drop of the suspension was deposited onto a perforated carbon membrane covered copper grid.

Cell cultures
HeLa cells (human tumor-derived epithelial cells) and RAW 264.7 cells (murine transformed macrophages) were purchased from ATCC (VA, USA). Both cell lines were cultured in RPMI 1640 medium supplemented with 10% heat inactivated FBS and 100 U ml⁻¹ gentamycin (with the addition of 50 μM β-mercaptoethanol and 20 mM HEPES for RAW 264.7 macrophages). Cells were maintained at 37 °C in humidified air containing 5% CO₂. Media and supplements were purchased from Lonza. When confluency reached 70%–80%, HeLa or RAW 264.7 cells were washed with phosphate-buffered saline (PBS), detached with trypsin or with SE buffer (PBS containing 2 mM EDTA and 2% FBS), respectively, and subcultured each 2–3 d. Before MLG exposure, cells were detached, counted and reseded in proper well size and density (see each particular experiment for details) and allowed to adhere overnight.

Flow cytometry
For cell viability experiments, HeLa or RAW 264.7 cells were seeded into 96-well culture plates at a density of 1 x 10⁵ cells per well and allowed to adhere overnight. Cells were exposed to different concentrations (ranging from 1 to 100 μg ml⁻¹) of MLG1 or MLG2 for 24 h. DMSO (20%) was used as positive control for cellular death. After incubation, RAW 264.7 supernatants were collected for further investigations while cells were harvested and stained with FITC-Annexin V (AnnV; BD Pharmingen 556419) and propidium iodide (PI, 0.2 μg ml⁻¹; Sigma-Aldrich) in a calcium containing buffer. The percentage of live (AnnV negative and PI negative), early apoptotic (AnnV positive and PI negative) and late apoptotic/necrotic (AnnV positive and PI positive) or late apoptotic/necrotic (AnnV positive and PI positive plus AnnV negative and PI positive) cells was determined by acquiring at least 50 000 events using a Gallios flow cytometer (Beckman Coulter, Villepinte, France) and analyzing the data with FlowJo software.

For the analysis of cell activation, after incubation with MLG1 or MLG2, RAW 264.7 cells were harvested and stained with PE-Rat anti-Mouse CD86 antibodies (Clone GL1, BD Pharmingen 553692). Lipopolysaccharide (LPS, 1 μg ml⁻¹) in combination with interferon γ (IFN γ, 1 ng ml⁻¹) was used as positive control. The macrophage-associated CD86 fluorescence intensity was determined by acquiring at least 50 000 events using the Gallios flow cytometer and analyzing the data on the live cell gated population with FlowJo software.

The flow cytometry results are a summary of the data from at least five separate experiments run in triplicate for each cell line. Data are presented ±SEM. Statistical analyzes were performed using a two-way ANOVA test followed by Bonferroni’s post-test. All p values <0.05 were considered significant.

ELISA
The concentration of interleukine 6 (IL6) and tumor necrosis factor alpha (TNF α) in RAW 264.7 cell supernatants were examined by a double sandwich ELISA. LPS (1 μg ml⁻¹) in combination with IFN γ (1 ng ml⁻¹) was used as positive control. Polyvinyl microtiter plates (Falcon) were coated overnight at 4 °C with 50 μl per well of purified Rat Anti-Mouse IL6 (BD Pharmingen 554400) or purified Hamster Anti-Mouse/Rat TNF α (BD Pharmingen 557516) antibodies diluted in 0.05 M carbonate buffer, pH 9.6. Non-specific sites were saturated with 100 μl per well of PBS containing 10% FBS for 1 h at 37 °C. 50 μl per well of culture supernatants or Recombinant Mouse IL6 (BD Pharmingen 554582) or Recombinant Mouse TNF (BD Pharmingen 554589) diluted in PBS-10% FBS, were added for 2 h at 37 °C. 50 μl per well of secondary Biotin Rat Anti-Mouse IL6 (BD Pharmingen 554402) or Biotin Rabbit Anti-Rat/Mouse TNF (BD Pharmingen 557432) diluted in PBS-10% FBS, were added for 1 h at 37 °C. 50 μl of streptavidin conjugated to horseradish peroxidase diluted in PBS 10% FBS were added per well. Plates were washed after each step with PBS containing 0.05% of tween20 (PBS-T). After 30 min incubation at 37 °C, plates were washed extensively and the enzymatic reaction was visualized by adding 75 μl per well of 3,3′,5,5′-tetramethylbenzidine diluted in 0.1 M citrate buffer (pH 5) in the presence of H₂O₂. The resulting absorbance was measured at 450 nm after the reaction was stopped with 25 μl per well of HCl 1 N. Statistical analyzes were performed as previously described in the flow cytometry section.

Xenopus breeding and maintenance
Eggs were obtained at the Ecolab laboratory resulting from the mating of two Xenopus adults following the hormonal injection of 50 IU of PMSG 500 (Pregnant Mare’s Serum Gonadotropin, Intervet, France) for the male and 750 IU of HCG (Human Chorionic Gonadotropin, Organon, France) for the female. Viable eggs were maintained in aquariums containing normal tap water filtered through active charcoal at
20 °C–22 °C until they reached the development stage appropriate for experimentation. The larvae were fed with dehydrated aquarium fish food every day (Tetraphyll®, Zolux, France).

**Xenopus exposure conditions**

The exposure was performed according to the International Standard 21427–1 [23] in semi-static exposure conditions with larvae at stage 50 [30]. The larvae were taken from the same laying to reduce inter-individual variability. They were exposed for 12 d in groups of 20 animals in crystallizing dishes containing either MLG concentrations (0, 0.1, 1, 10 and 50 mg l\(^{-1}\)) in reconstituted water (RW, i.e., chlorine-free water supplemented with nutritive salts, as described in ISO 21427–1 [294 mg l\(^{-1}\) CaCl\(_2\)·2H\(_2\)O, 123.25 mg l\(^{-1}\) MgSO\(_4\)·7H\(_2\)O, 64.75 mg l\(^{-1}\) NaHCO\(_3\), 5.75 mg l\(^{-1}\) KCl, pH 7) or control media, i.e. RW without MLG (negative control: NC and positive control: PC). The exposure was performed according to the International recommendation fascicle I [31]. Statistical analyzes were performed using SigmaStat 3.1 according to non parametric tests (Kruskal–Wallis followed by Dunn’s or Dunnet’s test and Mann–Whitney test) described in previous studies [24].

For the micronuclei test (MNT), at the end of exposure, a blood sample was obtained from each anaesthetized larva (MS222, Sandoz, France). Technical procedures are well described on the standardized recommendations [23]. The number of erythrocytes that contained one micronucleus or more (micronucleated erythrocytes, MNE) was determined in a total sample of 1000 erythrocytes per larva (MNE %). Based on median values and quartiles [32], the number of micronucleated erythrocytes per thousand, MNE % is presented with their 95% confidence limits expressed by the median ±1.57 × IQR inter-quartile range (upper quartile—lower quartile)/√n. The difference between the theoretical medians of the test groups and the theoretical median of the negative control group is significant to within 95% certainty if there is no overlap.

After blood puncturing, the general aspect of the larvae exposed to MLG was visually compared to that of NC group under the binocular. After dissection of some larvae of each group, their guts were then observed under the binocular (magnification ×15) to observe the presence or absence of MLG.

**Results and discussion**

**Toxicity on cellular models**

Inspecting nanomaterial acute toxicity on *in vitro* cellular models is an important first step for an appropriate assessment of their biocompatibility. For this purpose, tumor-derived epithelial cells (HeLa) were selected as valid human model to test the possible cytotoxicity of two MLG samples. It is important to point out that this cellular type represents a non-phagocytic model. The tests were performed also on a macrophage cell line (RAW 264.7). The latter was chosen as an important immune cell model, in order to compare the effects of MLG on the two cell lines and to further investigate the possible proinflammatory response triggered by MLGs. As a matter of fact, macrophages are one of the major actors during the primary immune response against infectious organisms and external materials, being able to phagocyte such entities and trigger secondary immune reactions [33]. The importance of MLG purification on cytotoxicity was also taken into account. For this purpose,
two different MLG samples were tested: MLG1, corresponding to the starting MLG batch and MLG2, consisting in MLG1 thermally treated to eliminate residual toluene molecules potentially adsorbed and entrapped between the graphene layers.

Cell viability was the first parameter that we analyzed. Both cellular models were exposed to increasing concentrations of MLG1 or MLG2 for 24 h. It is important to point out that the dispersion of both MLG samples in cell culture medium was not optimal. On the other hand, it was also possible to observe that MLG black spots often colocalized with the cells suggesting an interaction with the cell membranes. However, the general cellular morphology seemed comparable to control cells.

At the end of the incubation, the cell viability was determined by flow cytometry upon AnnV/PI staining. Data show that HeLa cells tolerated very well both MLG samples as no reduction of cell viability was observed (figure 2(A)). On the contrary, they were dramatically affected by control DMSO. A similar situation was monitored in

![Figure 1](image1.png)

Figure 1. Optical microscopy images of HeLa or RAW 264.7 cells after exposure to 100 μg ml⁻¹ of MLG1 or MLG2 for 16 h (B). Scale bar: 20 μm.

![Figure 2](image2.png)

Figure 2. Flow cytometry analysis of HeLa (A) and RAW 264.7 (B) cell viability exposed to different concentrations of MLG1 and MLG2 for 24 h. Two-ways ANOVA followed by Bonferroni’s post-test was performed to determine the statistical differences versus control cells and to compare the different graphene samples to each other (* p < 0.05; ** p < 0.01; *** p < 0.001).
the case of RAW 264.7 cells. In fact, MLG1 and MLG2 induced only a slight, but not significant, decrease of cell viability (of about 5%) in the macrophage cell line (figure 2[B]). No differences between the two MLG samples were evidenced.

Investigating the immunosafety of a new material is also of fundamental importance in the context of biomedical applications. In fact, the immune cells of the body may be activated by the external molecules and elicit a harmful inflammatory response [34]. To assess this second important parameter, RAW 264.7 macrophages were incubated with increasing concentrations of MLG1 or MLG2, as previously described, and the expression of the macrophage activation marker CD86 was analyzed by flow cytometry. The culture supernatants were also collected to evaluate the levels of two proinflammatory cytokines, (IL6 and TNF α) by ELISA. The results show that the levels of CD86 and cytokines were comparable to control cells, indicating that our MLG samples did not induce any proinflammatory response in RAW 264.7 macrophages (figure 3). On the contrary, cell activation and cytokine levels increased when the cells were treated with LPS/IFN-γ thus validating the test.

Our data on cell viability and cell activation suggest that MLG does not elicit any cytotoxic effects. However, such result is in contrast with some previous findings [22, 35]. To give a few examples, an early study reported the induction of high oxidative stress and caspase-3 activation (indicating apoptotic process) in neuronal rat cells exposed to pristine graphene (composed mainly of 3–5 layers) [36]. Another group showed that 3–4 layers GO can trigger cell cycle alterations, apoptosis, and oxidative stress in osteoblast and macrophage cell models [37]. However, graphene cytotoxicity depends on several factors such as graphene type (in terms of functionalities, number of layers and lateral size), graphene impurities and cell type tested, as it is reported in different reviews [22, 35]. Nevertheless, we can make a hypothesis to explain the discrepancy between our and literature data on graphene cytotoxicity. Our MLG samples are composed of several graphene sheets (2–20), while the majority of the works utilize single-layer or FLG (pristine or oxidized). Studies mainly focused on the graphene sheet lateral size, rather than the number of layers [38, 39]. The number of graphene layers is instead a very important parameter we have to consider because it determines specific surface area and bending stiffness [40]. In particular, the specific area is inversely proportional to the number of graphene layers. As a consequence, surface phenomena such as physical adsorption of molecules or catalytic chemical reactions, occurring within the biological environment, are reduced in our case. Since such events may be responsible for adverse biological reactions [40], this could explain the absence of cytotoxicity observed in our system. Furthermore, in our conditions, MLG dispersions were not optimal, as evidenced by the strong presence of aggregates (figure 1), thus decreasing even more the phenomena. Another hypothesis can also be made and concerns the material internalization. We assume that the stiffness of our MLG samples is substantially high [40]. Despite being phagocytic cells, macrophages may fail to internalize a material with this rigidity and even more large MLG aggregates, thus explaining the negligible effect on their cell viability and activation. The passive diffusion of the material through the plasma membrane, previously reported [39, 41], might also be negatively affected by MLG high stiffness. However, it has been reported that micro-sized graphene (lateral size comprised between 0.5 and 25 μm) composed of 4–25 layers, can be internalized by macrophages [41]. The authors hypothesize that the interaction between the hydrophobic basal surface of graphene microsheets with the inner hydrophobic region of the plasma membrane promoted the cellular uptake. Despite a lower lateral size, the aggregation status of our MLGs is likely sufficient to support the hypothesis of a lack of internalization. In the same
work, it has been also proposed that graphene microsheets, once internalized by macrophages, may be recognized as damage-associated molecular patterns that nonspecifically activate the innate immune response [42]. In our case, MLG hydrophobic surface area (average lateral size 2.3 μm and number of layers 2–20) is low and this represents another possible reason to explain the absence of proinflammatory reaction in RAW 264.7 macrophages. The importance of the material shape and size on macrophage inflammatory response was reported by us and other groups [38, 39]. Another previous study was focused on the cellular impact of GO sheets having different lateral sizes. The authors showed that only GO samples with sheets of about 2 μm triggered strong primary macrophage activation, contrarily to sheets of about 300 nm [38]. On the contrary, we have previously shown how, the smaller the GO lateral size, the higher the material internalization and the higher the proinflammatory effect was present, in both human and murine primary macrophages [39]. The cellular uptake of GO with the higher lateral size (1.32 μm in average) was less efficient compared to sheets with average lateral sizes of 0.27 and 0.13 μm, respectively. The large GO sample was also less cytotoxic than the two GOs in the nanoscale range towards macrophages. These previous results are fully consistent with the data obtained in this work, as our MLGs have an average lateral size of 2.3 μm.

These observations give a further demonstration that several parameters influence graphene toxicity. Despite we were not able to precisely evaluate the relative contribution of each parameter on MLG cytotoxicity, we can hypothesize that the relatively high number of layers and lateral size of MLGs combined with a certain degree of aggregation have a synergistic effect in determining the absence of toxicity in our cellular models. This is likely to be due to a poor material uptake and to the reduced hydrophobic surface area (related to the layer number of MLGs) which reduce surface adsorption phenomena considered responsible of promoting inflammatory responses [40].

Toxicity on Xenopus larvae

The highly pure MLG sample was further tested on X. laevis. In this work, the amphibian was used for a double scope. First, it is a valid model to further assess the in vivo impact of MLGs, giving more insights on the overall toxicity of the material. Second, the in vivo results give also information about the ecotoxicity of MLGs as X. laevis is a relevant environmental model of aquatic organism to study the ecotoxicity of nanoparticle [24–27].

The first parameter that we considered was larval survival after contact with the material, as index of acute toxicity. The results show no mortality in larvae exposed in presence of MLGs, whatever the concentration.

As second factor, we analyzed the variation of larval size, representing an indication of chronic toxicity, in response to MLGs. The measurements of the larval size show that larvae exposed in presence of 10 and 50 mg l⁻¹ of the material have significantly reduced size compared to the NC. However, no significant reduced size was observed at 0.1 and 1 mg l⁻¹ (figure 4).

The assessment of whether MLGs were able to induce damage in the genetic materials of living organisms, such as DNA mutations and/or chromosomal aberrations, is also very important. Micronucleus induction is a valid cytogenetic biomarkers used for measuring the genotoxic potential of agents. Figure 5 shows that larvae exposed to PC have significantly higher MNE % as compared to the NC group (17 ± 4.56 versus 2 ± 1.82), validating the MNT results. No genotoxicity via micronucleus induction in
erythrocytes of X. larvae was observed, whatever the concentration of MLGs. Indeed, the median values of MNE were $2 \pm 0.81$, $1 \pm 0.61$, $2 \pm 1.01$ and $2 \pm 1.22$ for respectively 0.1, 1, 10 and 50 mg l$^{-1}$.

X. larvae exposed during 12 d to MLGs in RW displayed a particular visual aspect compared to NC larvae (figure 6). Black agglomerates, supposed to be MLG, are increasingly observed in gills and intestines when the concentration of MLGs in media exposure increases. In addition, larval excretion of MLGs agglomerated with excrements is easily observed, for example in figure 7, after exposure to 50 mg l$^{-1}$ of MLGs.

Results on X. larvae show that MLG is substantially not toxic for this aquatic species. Such data are consistent with those found in the current literature on graphene ecotoxicity. Zanni et al showed that graphite nanoplates did not induce oxidative stress in the bacteria Pseudomonas aeruginosa, but 70% of bacterial viability was lost after 5 h of exposure to 250 mg l$^{-1}$ of the material [43]. They also showed no lifespan difference after exposing the nematode Caenorhabditis elegans to 250 mg l$^{-1}$ of nanoplates, although they observed the distribution of the material along the body of the nematode. In another study, the authors calculated the half maximal effective concentration (EC$_{50}$) values of 1.92 and 1.42 mg l$^{-1}$ after 30 min of exposure of the bacteria Vibrio fischeri to monolayer graphene and graphene nanopowder, respectively [44]. They also calculated EC$_{50}$ of 1.14 and 2.25 mg l$^{-1}$ after 72 h of exposure of the algae Dunaliella tertiolecta to monolayer graphene and graphene nanopowder, respectively. No acute toxicity against the crustacean Artemia salina was detected although aggregation of graphene into the gut was evidenced [44]. Guo and collaborators showed $^{14}$C-labeled graphene content in the body of the crustacean Daphnia magna (1% of the organism dry mass) after 24 h of exposure to 250 $\mu$g l$^{-1}$ of the material [45]. Another work reported the distribution of 100 mg l$^{-1}$ of multi-functional graphene from head to tail of the embryos of the fish Danio rerio. However, the material did not induce abnormalities in the organisms [46].

In our case, only the X. larvae growth rate seemed to be affected by the material to some extent. Indeed, the larval size was decreased by about 40% only upon exposure to the highest concentration of MLGs (50 mg l$^{-1}$). This profile response is usually observed in X. larvae exposed under the same conditions to carbon nanotubes [24, 25]. Several hypotheses have already been proposed to explain this phenomenon. For instance, carbon-based nanomaterial uptake may lead to digestive (gut) and respiratory (gills) obstruction causing some exchange gas dysfunctions. A
competition between nutrients and carbon based nanomaterials might be another explanation. Carbon-based nanomaterials, quickly sediment into the water column at the bottom of experimental dishes forming agglomerates with food and excreta during the exposure. When larvae browse at the bottom of the experimental dishes, massive larval uptake of the agglomerates occurs, thus leading to respiratory and/or intestinal clogging.

On the other hand, MLG did not induce larval mortality and genotoxicity, at any of the concentrations tested. Such data support the results obtained on the human epithelial cells (HeLa) and on murine macrophages (RAW 264.7). MLG agglomerates could be then excreted without triggering any genetic toxicity towards the amphibian. One can speculate that, also in vivo, our particular MLGs cannot be internalized by the cells they encounter. Such cells are mainly represented by the epithelial cells of the respiratory and digestive tracts, which are the main way of entry in our case. Consequently, MLG may not be able to reach other body compartments such as the blood flow, thus explaining also the absence of genotoxicity towards erythrocytes. In fact, our previous studies using transmission electron microscopy, Raman spectroscopy and histology could not clearly evidence the translocation of carbon nanotubes through the intestinal barrier in Xenopus [24, 47].

To conclude, in this multifaceted work, MLGs composed of 2–20 graphene layers, were tested on two mammalian cell models and on X. laevis as an important in vivo and environmental model organism. MLG showed to be substantially not toxic towards our cellular models and X. larvae. We hypothesize that the absence of harmful effects both in the in vitro and in vivo models can be ascribed to different factors. The first reason might be represented by the failure in MLG cellular internalization, due to the particular size-related characteristics of our material [40], and to its poor dispersibility in aqueous media. It is conceivable that cells may encounter more difficulty in internalizing a material aggregated by endocytosis/phagocytosis, thus explaining the absence of toxicity. On the other hand, the specific surface area of our MLG is low and less exposed to the surrounding biological environment, compared to monolayer or FLG-based materials [40]. For this reason, the material might not be recognized by macrophages as a hydrophobic damage-associated molecular pattern, able to activate the innate immune response [42]. In addition, surface phenomena such as physical adsorption of molecules or catalytic chemical reactions, occurring within the biological environment, are reduced in our case. Since such events may be responsible for adverse biological reactions [40], this could also explain the absence of cytotoxicity observed in our system.

This study gives an important contribution on the impact of MLG on health and environment and further evidences the importance of graphene size, shape and dispersibility on its biological impact and on the synergistic effects that different parameters might have in triggering toxic effects.

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