In vivo characterization of carbon dots–bone interactions: toward the development of bone-specific nanocarriers for drug delivery

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ABSTRACT
Current treatments for osteoporosis and other bone degenerative diseases predominately rely on preventing further bone erosion rather than restoring bone mass, as the latter treatments can unintentionally trigger cancer development by indiscriminately promoting cell proliferation. One approach to circumvent this problem is through the development of novel chemical carriers to deliver drug agents specifically to bones. We have recently shown that carbon nanodots (C-dots) synthesized from carbon nanopowder can bind with high affinity and specificity to developing bones in the larval zebrafish. Larval bones, however, are physiologically different from adult bones in their growth, repair, and regeneration properties. Here we report that C-dots can bind to adult zebrafish bones and that this binding is highly specific to areas of appositional growth. C-dots deposition occurred within 30 minutes after delivery and was highly selective, with bones undergoing regeneration and repair showing higher levels of C-dots deposition than bones undergoing normal homeostatic turnover. Importantly, C-dots deposition did not interfere with bone regeneration or the animal’s health. Together, our results establish C-dots as a potential novel vehicle for the targeted delivery of drugs to treat adult bone disease.

Introduction
Bone diseases such as osteoporosis affect over 54 million individuals annually in the US alone (Wright et al., 2014), imposing a heavy burden to society that could be mitigated with improved diagnostic techniques and expanded treatment options. For osteoporosis, early diagnosis is key for the most successful treatment prognosis, as current treatments predominantly rely on preventing further bone erosion and not in restoring bone mass, as drugs that promote bone growth can lead to cell proliferation in other tissues and increase a patient’s risk to cancer (Marie & Kassem, 2011). Bone disease diagnostics rely on X-ray imaging methods and MRI or CT scans, with novel, higher sensitivity, fluorescent-based technologies currently under development (Chen et al., 2014; Gruneboom et al., 2019; Jung et al., 2019; Yan et al., 2019). The development of more sensitive diagnostic tools for early bone-loss detection and novel treatment methods for stimulating bone growth without affecting other tissues will significantly ameliorate the societal and personal cost of bone diseases.

Carbon-based nanoparticles or carbon dots (C-dots) have emerged as novel therapeutic and diagnostic biomaterials due to their unique, tunable physicochemical properties (Zhou, Mintz, et al., 2019; Dan et al., 2020). Common properties shared by all C-dots include small size (<10 nm), high carbon content, robust photostability, and bright fluorescence (Mintz et al., 2019; Zhou, Mintz, et al., 2019). While several physicochemical factors contribute to these properties, the most influential are core configuration and surface functional groups (Mintz et al., 2019; Yan et al., 2019; Zhou, Zahran, et al., 2019). Importantly, the functional groups at the surface can be modified during their synthesis to fine-tune the C-dots’ photoluminescent properties (Zhou, Mintz, et al., 2019; Dan et al., 2020), or to adapt C-dots to function as pharmaceutical nanocarriers (Zheng et al., 2014; Zhou, Liyanage, et al., 2019). For example, the surface of C-dots can be passivated with polyethylene glycol (Sun et al., 2006; Peng et al., 2020), and then conjugated covalently or noncovalently with diverse therapeutic agents drugs (Thakur et al., 2014; Iannazzo et al., 2017; Hettiarachchi et al., 2019; Liyanage et al., 2020). These modifications, however, can have unexpected consequences for both the C-dot vehicle and the drug cargo, including affecting the solubility, therapeutic efficacy, and systemic clearance of the drug (Misra et al., 2015; Li, Amat, et al., 2016; Li et al., 2020). To fully exploit...
C-dots’ theragnostic potential, it is imperative to first characterize unmodified C-dots.

We recently described that C-dots synthesized from carbon nanopowder have a high affinity and specificity for developing zebrafish bones (Li, Skromne, et al., 2016; Peng et al., 2017). When carbon nanopowder-derived C-dots were injected into 5-day-old zebrafish larvae, their deposition was observed in opercular (intramembranous) and vertebrae (endochondral) bones, but not in non-skeletal tissues (Li, Skromne, et al., 2016; Peng et al., 2017). C-dots deposition on bones was dependent on mineralization, as manipulations promoting or interfering with bone mineralization increased or decreased C-dot deposition respectively (Li, Skromne, et al., 2016). Of note, C-dots binding to bones was only observed when the C-dots were synthesized from carbon nanopowder through oxidation in hydrothermal conditions, and not when synthesized from citric acid or EDA using solvo-thermal methods (Peng et al., 2017). This difference in binding may be due to the presence of different chemical groups at the surface of the C-dots: while nanopowder-derived C-dots are rich in negatively charged carboxyl groups that can interact with positive calcium ions in the bone’s matrix, the surface of citric acid- and EDA-derived C-dots is rich in positive amino groups (Peng et al., 2017). These results suggest that the unique physicochemical properties of carbon nanopowder C-dots are critical for their binding to the larval mineralized bone.

Zebrafish provides a robust, in vivo model to test C-dots’ interaction with biological tissues, as their transparency allows direct observation of C-dots’ photoluminescence. Importantly, bone formation and maintenance in zebrafish are remarkably similar to that of other vertebrates (Witten et al., 2017; Busse et al., 2020; Tonelli et al., 2020). During embryogenesis in fish, birds, and mammals, bones develop either through the direct aggregation of bone-forming cells (intramembranous ossification; e.g. cranial bones) or through the deposition of a mineral matrix on a collagen scaffold (endochondral ossification; e.g. long bones) (Salhotra et al., 2020). Once formed, adult bones undergo homeostatic turnover and can continue to increase in diameter through the process of appositional growth, whereby new mineralized tissue is added to the bone’s surface (Rauch, 2005; Suniaga et al., 2018; Salhotra et al., 2020). These appositional growth and remodeling processes are carried out in zebrafish by similar skeletal cells and ossification mechanisms that have been observed in mammals (Busse et al., 2020; Tonelli et al., 2020). In addition, zebrafish can rapidly regenerate caudal fin tissue, including bony fin-rays, in less than 6 days (Busse et al., 2020; Tonelli et al., 2020). This remarkable regenerative capacity, together with transparency and thinness, makes the caudal fin of the adult zebrafish an excellent model to study bone homeostasis, repair, and regeneration.

Here, we investigated the interactions between carbon nanopowder C-dots and skeletal tissues of adult zebrafish, reporting that C-dots deposit on adult bones undergoing homeostatic turnover, appositional growth, and regeneration. This is an important first step toward developing C-dots as theragnostic agents for the treatment of adult bone diseases.

We report that, irrespectively of delivery method, C-dots bind with high affinity and specificity to mineralized adult bones, particularly those undergoing rapid growth. C-dot deposition occurred within the first 30 minutes after injection and was found to co-localize to Alizarin Red Complexone (ARC), a well-known histological marker of areas of bone growth (Hoyte, 1960). Importantly, C-dots deposition to bones did not interfere with normal bone physiological processes and growth, or affect animal’s viability. Taken together, our results highlight the vast potential of photoluminescent C-dots as a tool for the diagnosis and treatment of adult bone diseases.

Materials and methods

Carbon dot synthesis and analysis

Carbon nanodots (C-dots) were synthesized from carbon nanopowder and purified using our previously reported procedure (Li et al., 2015; Li, Skromne, et al., 2016). To obtain C-dots in powder form, the C-dot solution was lyophilized in a Rotovap evaporator at 60°C. The morphology and photoluminescent properties of as-prepared C-dots were confirmed using transmitted electron and epifluorescence microscopy, as previously described (Li et al., 2015; Li, Skromne, et al., 2016).

Zebrafish care, tail amputation, injection, and bone staining

Wild type (TAB-5; (LaFave et al., 2014)) and Casper (mpv1779; mitfam2; (White et al., 2008)) zebrafish were obtained from the Zebrafish International Resource Center (Eugene, OR) and maintained at the University of Richmond animal facility following standard husbandry protocols (Westerfield, 1995). All protocols and procedures were reviewed and approved by the Institutional Animal Care and Use Committee. All experiments were performed on animals 4–6 months of age. For amputations, fish were fully anesthetized in 0.2 mg/ml Tricaine (pH 7.0) until unresponsive to touch, placed on an inverted glass Petri dish covered with Parafilm, and a sterile blade was used to cut 40 mm away from the distal tip of the caudal fin. For injection, fish were anesthetized and weighed to standardize the mass of C-dots injected per body weight (μg/g). Then, fish were positioned on a wet sponge under the microscope and injected intraperitoneally or intravascularly with C-dots using a 36G Nanofill microsyringe attached to an electronically controlled micropump (UMP3 UltraMicroPump, WPI), as previously described (Pugach et al., 2009; Kinkel et al., 2010). After manipulations, fish were allowed to recover for 30 min and returned to the animal facility where they received standard care for the duration of the experiment.

Live staining of mineralized structures was done using Alizarin Complexone (ALC; Sigma-Aldrich, A-3882). First, fish were sedated in Tricaine solution in fish facility water (0.1 mg/ml; pH 7.0). Then, fish were stained in a solution of ALC (10 mg/ml) and Tricaine (0.01 mg/ml; pH 7.0), for 30 min. To remove excess dye following exposure, fish were quickly
rinsed in three sequential washes of fresh facility water and allowed to recover from sedation for 60 min before collecting the caudal fins for histology. Following fin collection, fish were allowed to recover as above.

**Histology**

For histological sectioning, fish were anesthetized and the regenerated fin was amputated at a site proximal to the original cut. For unmanipulated controls, the cut was done 40 mm away from the distal tip of the caudal fin. Immediately after caudal fin collection, the fish was placed in a recovery tank and the fin was splayed out by floating it on a large drop of 4% paraformaldehyde in 1x PBS inside a Petri dish. After a few min, once the fin was submerged in the droplet, the Petri dish was closed and the fin was fixed for 4 h. The fixative was then removed and replaced with a 10% sucrose solution. The Petri dish was sealed and stored at 4°C overnight. Samples were processed within 5 days. Fins were embedded in Clear Frozen Section Compound (VWR), frozen, and sectioned at 5 microns in a Leica CM1520 cryostat. Sections were collected on positively charged slides (Globe Scientific), allowed to dry for 30 min, and covered with mounting media (30% glycerol in 1x PBS, 5 mg/ml n-Propyl gallate, 2.5 μg/ml DAPI). Slides were kept in the dark at 4°C and sections were imaged within one week.

**Imaging and analysis**

For imaging, zebrafish were anesthetized and placed on a glass Petri dish with the caudal tail splayed to separate the fin rays. Whole fin images were acquired under brightfield and epi-fluorescent light using a Zeiss Discovery V.20 dissecting microscope and a Zeiss Axiocam MRc camera. Filter sets for Fluorescein (green; 488/525 nm) and Texas Red (red; 596/615 nm) were used to detect C-dots and ALC, respectively. For consistency across experiments and biological replicates, all fluorescent images were taken using time exposures of 1 and 2.5 s. Images were processed using Zeiss AxioVision SE64 v4.9.1 and fluorescence quantification was done using ImageJ. Signal intensity in regions of interest or along the profile line was recorded and analyzed in Excel. Graphical visualization of data was done in Excel or RStudio. The RStudio code used for fluorescence analysis is available as Supplemental Material on the publisher’s webpage.

**Results and discussion**

**C-dots bind to adult bones undergoing homeostatic turnover, repair, and regeneration**

To gain insight into the adult theragnostic potential of carbon nanopowder-derived C-dots, we analyzed their toxicity, bone binding dynamics, and photoluminescence in transparent adult zebrafish (Casper strain (White et al., 2008)). We analyzed C-dots’ bone deposition using standard 488/525 nm excitation/emission Fluorescein filter set to capture their intrinsic photoluminescence (excitation peaks between 360–540 nm and emission between 500–600 nm (Li, Amat, et al., 2016)). Three different delivery methods were used to determine the ability of C-dots to bind adult bones, one cutaneous and two through injection (Figure 1). To determine if C-dots can be adsorbed cutaneously, we applied a concentrated solution of C-dots to the caudal fin or the flank of the fish. Neither application resulted in the labeling of local bones (Figure 1(A,A’,B,B’)) and data not shown). Similarly, topical application of C-dots to a healed caudal fin wound undergoing regeneration did not label local bones (Figure 1(C,C’,D,D’)). In contrast, C-dots applied to a wound with exposed bones labeled the exposed tips of the bones (Figure 1(E,E’)). Notably, none of the bone tissues that regenerated following C-dots exposure were labeled, indicating that following deposition, C-dot binding became fixed (Figure 1(F,F’)). We next employed two injection delivery methods, intraperitoneal and intravascular (retro-orbital). Delivery of C-dots (20 μg/g fish) through either method effectively labeled intact cranial, ribs, and fin bones (Figure 1(G–O,G’–O’)). This labeling was most apparent in the fish fins because of the tissue’s reduced thickness and isolated position (Figure 1(G–O,G’–O’)). These results indicate that direct wound application and injection methods are effective for delivering C-dots into adult fish, with intravascular injections resulting in the most consistent deposition of C-dots on bones (not shown). Together with our previous results in larvae (Li, Amat, et al., 2016), this suggests that C-dots can bind with high specificity to skeletal elements irrespective of bone age.

To explore the binding of C-dots to wounded and regenerating adult bones, we administered C-dots to adult zebrafish that had undergone amputation of the caudal fin and were at different stages of regeneration. This approach allowed us to examine the binding of C-dots to fin ray bones undergoing normal homeostatic turnover as well as regenerative growth. When C-dots were delivered prior to the initiation of fin bone regeneration at 1 day post-amputation (dpa; Figure 2(A)), regenerated bones examined at 8 dpa...
were not labeled (Figure 2B–E). However, when C-dots were delivered during active fin bone formation (e.g. 4 dpa; Figure 2F), regenerated bones were strongly photoluminescent (Figure 2G–J). The observation that C-dots delivery at 4 but not 1 dpa labeled regenerating bones at 8 dpa suggests that C-dots were quickly cleared from circulation. Attempts to determine if C-dots were metabolized or excreted were unsuccessful, as C-dot concentrations were below the detection limit of our assays (ELISA and TEM; data not shown). Therefore, at this moment, we cannot rule out the possibility that C-dots were removed from the circulatory system through mechanisms other than bone deposition. Nonetheless, once in bones, C-dots label could be detected for at least eight weeks (data not shown), suggesting that binding to adult bones may be permanent.

**C-dots deposition on regenerating bone is dose-dependent, swift, and homogeneous**

The photoluminescence intensity of C-dots bound to regenerating bones was qualitatively more intense and homogeneous than the binding to non-regenerating bones, providing an assay to study C-dots binding to ossifying tissue. To begin testing the magnitude of C-dot binding to regenerating bone as a function of C-dots dosage, we repeated the amputation experiment followed by the intravascular administration of increasing amounts of C-dots (Figure 2F). The amount of C-dots in solution delivered to each fish was normalized to the weight of the fish (µg/g). The lowest amount at which we were able to detect C-dots binding to regenerating bones was 5 µg/g (Figure 2K). The photoluminescence intensity increased until 40 µg/g, the highest dose tested (Figure 2K). Higher concentrations were not tested because concentrations above 40 µg/g clogged the microneedles used for injections and the use of larger needles damaged the fish’s vasculature at the site of injection. To quantify C-dots’ binding, we measured the photoluminescence intensity of a 100 µm square area located 100 µm away from the amputation site in regenerating fin rays 4, 6, 8, and 10 (ventral to dorsal position). These values were averaged and normalized to the intrinsic background of corresponding fish as well as to control fish (0 µg/g). The normalized averages were then plotted against C-dots’ dosage to determine the magnitude of deposition. Based on three independent experiments, there is a linear relationship between C-dots dosage and photoluminescence intensity (Figure 2L): linear regression, $R^2 > 0.96$; Standard t-test, $df = 3, p < 0.002$). Trendlines did not plateau, suggesting that tissue saturation was not achieved. Together, these results suggest that concentrations of C-dots ranging from 5–40 µg/g can label regenerating bone without compromising the fish’s health or the process of bone regeneration. Due to concern...
that large volumes of C-dots may cause tissue damage during injection, we limited the concentration of C-dots to 20 \( \mu \text{g/g} \) in subsequent experiments.

Next, to determine whether C-dots deposition on bones is dependent on the state of regeneration, we repeated the experiment varying the time of injection after amputation. The time of C-dot injection varied from 2 to 7 dpa, and all the regenerated fins were imaged at 8 dpa (Figure 3(A)). We then quantified the photoluminescence intensity profile of the C-dots deposited in the fourth most ventral regenerated fin ray (Figure 3(B)). Compared to controls, injection at 2 dpa resulted in the least amount of C-dots deposition, which was primarily restricted to the site of amputation. Injection at subsequent days increased the area of C-dots deposition along regenerating bones, with the whole regenerated ray being labeled at 7 dpa (Figure 3(A)). Signal quantification of the labeled bones revealed an inverse correlation between the amount of tissue labeled and the photoluminescence intensity of the labeling: the more area labeled, the lower the intensity of the photoluminescent signal (Figure 3(B)). These observations suggest that C-dots deposition after fin amputation occurs at all stages of bone regeneration, resulting in a homogeneous distribution of fluorescent labeling throughout the bone tissue distal to the amputation site. Clearance of C-dots from circulation was swift, as C-dots deposition occurred in a regenerated bone already present in the fin, and not in bone regenerated in subsequent days. For example, injection at 3 dpa only labeled the bone that had regenerated at that point, and not the bone that regenerated between 4–7 dpa (Figure 3). Together these results suggest that C-dots have a short circulatory half-life, depositing in available bones at the time of injection.

**C-dots deposit at areas of appositional bone growth**

To investigate the dynamics of C-dot deposition at the tissue level, we analyzed the distribution of C-dots’
photoluminescence in transverse sections of regenerating caudal fins. Adult zebrafish were injected with C-dots 1, 4 or 8 dpa and, after 1-h of labeling, the fin was harvested, cryosectioned, and imaged using confocal microscopy. We did not observe any histological differences in the regenerated bones between control and C-dot injected fish (Figure 4 and data not shown), supporting the gross morphological observations that C-dots deposition does not interfere with bone regeneration processes. At all stages of regeneration, bones proximal to the amputation site were thicker, more mineralized, and exhibited a strong C-dot signal (Figure 4(B)). To test if bone age is a determinant of C-dot deposition, we analyzed cross-sections of bones at different stages of regeneration at comparable proximal-distal positions. Comparison of representative sections adjacent to the amputation site revealed weak labeling of newly regenerated bones at 1 dpa, strong and even labeling at 4 dpa, and strong labeling of bone surfaces at 8 dpa (Figure 4). Given that the diameter of...
bones increases by the repeated addition of ossified tissue at the surface (appositional growth (Rauch, 2005)), our observations suggest that C-dot deposition occurs in areas of bone growth.

To directly test C-dots deposition in areas of tissue growth, we examined the distribution of C-dots relative to that of the dye Alizarin Red Complexone (ARC). Intravital staining of bones by Alizarin red dyes is a well-established histological method used in mice to distinguish areas of active bone growth, as the dye is preferentially taken up by these areas compared to regions where bone growth has slowed down or ceased (Hoyte, 1960). Thus, co-localization of C-dots and ALC signals would indicate that C-dots preferentially bind to areas of active bone growth. To validate the use of ALC in zebrafish, we exposed fish undergoing caudal fin regeneration to ALC at 8 dpa. In these fish, regenerated fin bones were strongly stained with ALC (Figure 5(A)). In cross-sections, ALC's stain distribution was homogeneous in bones closer to the fin tips (Figure 5(D,D')), and superficial in bones closer to the amputation site (Figure 5(E,E')). These findings are similar to those observed during the normal growth of bones in mammals (Hoyte, 1960), supporting the use of ALC as a viable method for identifying areas of bone growth in zebrafish.

Next, we investigated the relationship between C-dot deposition and areas of ALC staining. This analysis, however, had to be done indirectly, as C-dots’ 500–600 nm emission range overlaps with the emission peak of ALC at 580 nm. Using an exclusion approach, we could identify areas of growth that were ALC-positive and C-dots-negative by looking for 580 nm fluorescence in the absence of 525 fluorescence. We began this analysis by injecting fish with C-dots at 7 dpa and staining them with ALC one day later (Figure 5(B)). Because bone layer formation takes more than 24 h, we expected to see extensive C-dots and ALC signal co-localization. In all sections examined, the 525 and 580 nm signals co-localized (Figure 5(F,F',G,G')). At the distal tips of the fins, signal co-localization in newly formed bones were observed at the core and surface of the tissue (Figure 5(F,F')), whereas closer to the amputation site, signal co-localization in

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![Diagram](image.png)

**Figure 5.** C-dots deposit at sites of active bone mineralization. (A–C) Regenerated caudal fin of uninjected fish (A), injected with C-dots at 7 dpa (B), or injected at 4 dpa (C). All fish were stained with Alizarin Complexone (ALC) at 8 dpa and imaged. Composite images were generated by superimposing photographs obtained using transmitted light, a 488/525 nm filter set to capture C-dots’ fluorescence (pseudo colored green), and a 596/615 nm filter set to capture C-dots/ALC’s fluorescence (pseudo colored magenta). ALC-positive areas (indictive of ossification) that lack C-dots appear magenta, and areas of ossification with C-dot deposition appear white. Posterior end of fin toward top of images. Site of amputation is indicated with arrowheads and the plane of sections (D–L) are indicated with dotted lines. (D–L) Transverse sections of fins, imaged and pseudo colored as in (A)–(C). (D'–L') Magnified bone regions from D–E (white squares). Areas of C-dot deposition are indicated with white arrows. Ossified regions devoid of C-dots are indicated with black arrows. Asterisks indicate the bone matrix. Double asterisk in K' indicates areas of matrix buildup at the site of amputation. Scale bars are 500 μm for (A)–(C), 50 μm for (D)–(L) and 5 μm for (D')–(L').
older bones was observed only at the tissue’s surface (Figure 5(G,G)). This signal co-localization suggested that C-dots deposition occurred in areas of appositional growth. To further test this idea, we spatially separated temporal areas of growth by injecting fish with C-dots at 4 dpa and staining them with ALC four days later. In this experiment, we hypothesized that areas of early bone growth would fluoresce at 525 and 580 nm indicating C-dots deposits, whereas areas of late bone growth would fluoresce only at 580 nm indicating ALC staining. Consistent with this hypothesis, whole-mount analysis of the fins revealed that regenerated bones had an even distribution of the 580 nm fluorescent signal and a proximally-confined distribution of the 525 nm fluorescent signal (Figure 5(C)). Sectioning of the tissue further confirmed our hypothesis. Bones in distal regions of the fin showed 580 nm fluorescence only at their surface, indicating that areas of new growth stained only with ALC (Figure 5(H,H')). In contrast, bones at the amputation site or that did not undergo regeneration were positive for both fluorescent signals at the bone surface, indicating C-dots deposits and, presumably, ALC staining (Figure 5(K,K',L,L')). Strikingly, between these two regions, bones that were in the process of regeneration at the time of C-dot injection showed a dual staining pattern: the core portion of the bone was positive for both fluorescent signals indicative of C-dot deposits, while the surface of the bone was only positive for the 580 nm signal indicative of ALC staining (Figure 5(I,I',J,J')). We interpret this dual staining pattern to indicate that, as regenerating bones increase in size due to the apposition of new layers of tissue at their surface, early deposits of C-dots become surrounded by new tissue that stains positive for ALC. This observation, together with our previous result showing that C-dots and ALC signals overlap when treatments are done within 24 h of each other suggests that C-dots bind to areas of bone growth.

Conclusions
The discovery that C-dots bind to areas of adult bone growth without interfering with the biological processes of homeostatic turnover, repair, and regeneration opens the possibility for their use as theragnostic agents for diseases such as osteoporosis. Current osteoporosis treatments rely on early diagnosis to mitigate further bone erosion, as limited treatments exist to restore lost bone mass. C-dots’ intrinsic photoluminescence and bone-binding properties could revolutionize osteoporosis diagnostic tools by allowing clear visualization of the active and inactive areas of bone growth. For treatment, C-dots could be used as fluorescently traceable, bone-specific, drug-delivery agents. We have previously shown that surface modification of C-dots with ethylenediamine or glutamic acid does not result in loss of fluorescence or bone binding properties in larvae, providing proof of principle of their possible use as a bone-specific drug delivery method (Peng et al., 2017). Future work is needed to establish that drug conjugation to C-dots does not result in loss of the drug’s biological activity. Together, our results provide an important foundation for future applications of C-dots as a drug delivery vehicle for the treatment of adult skeletal diseases and traumatic bone injuries.

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Disclosure of interests
No potential conflict of interest was reported by the author(s).

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References
Busse B, Galloway JL, Gray RS, et al. (2020). Zebrafish: an emerging model for orthopedic research. J Orthop Res 38:925–36.
Chen G, Qiu H, Prasad PN, Chen X. (2014). Upconversion nanoparticles: design, nanochemistry, and applications in theranostics. Chem Rev 114:3161–214.
Dan Q, Xiayan W, Yuping B, Zaicheng S. (2020). Recent advance of carbon dots in bio-related applications. J Phys Mater 3:022003.
Gruneboom A, Kling L, Christiansen S, et al. (2019). Next-generation imaging of the skeletal system and its blood supply. Nat Rev Rheumatol 15:533–49.
Hettiarachchi SD, Graham RM, Mintz KJ, et al. (2019). Triple conjugated carbon dots as a nano-drug delivery model for glioblastoma brain tumors. Nanoscale 11:6192–205.
Hoyte DAN. (1960). Alizarin as an indicator of bone growth. J Anat 94: 432–42.
Iannazzo D, Ziccarelli I, Pistone A. (2017). Graphene quantum dots: multifunctional nanoplatforms for anticancer therapy. J Mater Chem B 5: 6471–89.
Jung JS, Jo D, Jo G, Hyun H. (2019). Near-infrared contrast agents for bone-targeted imaging. Tissue Eng Regen Med 16:443–50.
Kinkel MD, Eames SC, Philipson LH, Prince VE. (2010). Intrapeloneal injection into adult zebrafish. J Vis Exp 42:2126.
LaFave MC, Varshney GK, Vemulapalli M, et al. (2014). A defined zebrafish line for high-throughput genetics and genomics: NHGRI-1. Genetics 198:167–70.
Li S, Amat D, Peng Z, et al. (2016). Transferrin conjugated nontoxic carbon dots for doxorubicin delivery to target pediatric brain tumor cells. Nanoscale 8:16662–9.
Li S, Skromne I, Peng Z, et al. (2016). “Dark” carbon dots specifically “light-up” calcified zebrafish bones. J Mater Chem B 4:7398–405.
Li SH, Wang LY, Chusuei CC, et al. (2015). Nontoxic carbon dots potently inhibit human insulin fibillation. Chem Mater 27:1764–71.
Li X, Vinothini K, Ramesh T, et al. (2020). Combined photodynamic-chemotherapy investigation of cancer cells using carbon quantum dot-based drug carrier system. Drug Deliv 27:791–804.
Liyanage PY, Zhou Y, Al-Youbi AO, et al. (2020). Pediatric glioblastoma target-specific efficient delivery of gemcitabine across the blood-brain barrier via carbon nitride dots. Nanoscale 12:7927–38.
Marie Pj, Kassem M. (2011). Osteoblasts in osteoporosis: past, emerging, and future anabolic targets. Eur J Endocrinol 165:1–10.
Mintz KJ, Zhou Y, Leblanc RM. (2019). Recent development of carbon quantum dots regarding their optical properties, photoluminescence mechanism, and core structure. Nanoscale 11:4634–52.

Misra SK, Ohoka A, Kolmodin NJ, Pan D. (2015). Next generation carbon nanoparticles for efficient gene therapy. Mol Pharm 12:375–85.

Peng Z, Ji C, Zhou Y, et al. (2020). Polyethylene glycol (PEG) derived carbon dots: preparation and applications. Appl Mater Today 20:100677.

Peng Z, Mianji EH, Zhou Y, et al. (2017). Carbon dots: promising biomaterials for bone-specific imaging and drug delivery. Nanoscale 9:17533–43.

Pugach EK, Li P, White R, Zon L. (2009). Retro-orbital injection in adult zebrafish. J Vis Exp 34:1645.

Rauch F. (2005). Bone growth in length and width: the Yin and Yang of bone stability. J Musculoskelet Neuronal Interact 5:194–201.

Salhotra A, Shah HN, Levi B, Longaker MT. (2020). Mechanisms of bone development and repair. Nat Rev Mol Cell Biol 21:696–711.

Sun YP, Zhou B, Lin Y, et al. (2006). Quantum-sized carbon dots for bright and colorful photoluminescence. J Am Chem Soc 128:7756–7.

Suniaga S, Rolvien T, Vom Scheidt A, et al. (2018). Increased mechanical loading through controlled swimming exercise induces bone formation and mineralization in adult zebrafish. Sci Rep 8:3646.

Thakur M, Pandey S, Mewada A, et al. (2014). Antibiotic conjugated fluorescent carbon dots as a theranostic agent for controlled drug release, bioimaging, and enhanced antimicrobial activity. J Drug Deliv 2014:282193.

Tonelli F, Bek JW, Besio R, et al. (2020). Zebrafish: a resourceful vertebrate model to investigate skeletal disorders. Front Endocrinol 11:489.

Westfield M. (1995). The zebrafish book: A guide for the laboratory use of zebrafish (Brachydanio rerio). Eugene, OR: University of Oregon Press.

White RM, Sessa A, Burke C, et al. (2008). Transparent adult zebrafish as a tool for in vivo transplantation analysis. Cell Stem Cell 2:183–9.

Witten PE, Harris MP, Huysseune A, Winkler C. (2017). Small teleost fish provide new insights into human skeletal diseases. Methods Cell Biol 138:321–46.

Wright NC, Looker AC, Saag KG, et al. (2014). The recent prevalence of osteoporosis and low bone mass in the United States based on bone mineral density at the femoral neck or lumbar spine. J Bone Miner Res 29:2520–6.

Yan F, Sun Z, Zhang H, et al. (2019). The fluorescence mechanism of carbon dots, and methods for tuning their emission color: a review. Mikrochim Acta 186:583.

Zheng M, Liu S, Li J, et al. (2014). Integrating oxaliplatin with highly luminescent carbon dots: an unprecedented theranostic agent for personalized medicine. Adv Mater 26:3554–60.

Zhou Y, Liyanage PY, Devadoss D, et al. (2019). Nontoxic amphiphilic carbon dots as promising drug nanocarriers across the blood-brain barrier and inhibitors of beta-amyloid. Nanoscale 11:22387–97.

Zhou Y, Mintz KJ, Sharma SK, Leblanc RM. (2019). Carbon dots: diverse preparation, application, and perspective in surface chemistry. Langmuir 35:9115–32.

Zhou Y, Zahrani EM, Quiroga BA, et al. (2019). Size-dependent photocatalytic activity of carbon dots with surface-state determined photoluminescence. Appl Catal B 248:157–66.