Serotonin: a novel bone mass controller may have implications for alveolar bone

Carlo Galli1,2, Guido Macaluso*1 and Giovanni Passeri2

Abstract
As recent studies highlight the importance of alternative mechanisms in the control of bone turnover, new therapeutic approaches can be envisaged for bone diseases and periodontitis-induced bone loss. Recently, it has been shown that Fluoxetine and Venlafaxine, serotonin re-uptake inhibitors commonly used as antidepressants, can positively or negatively affect bone loss in rat models of induced periodontitis. Serotonin is a neurotransmitter that can be found within specific nuclei of the central nervous system, but can also be produced in the gut and be sequestered inside platelet granules. Although it is known to be mainly involved in the control of mood, sleep, and intestinal physiology, recent evidence has pointed at far reaching effects on bone metabolism, as a mediator of the effects of Lrp5, a membrane receptor commonly associated with Wnt canonical signaling and osteoblast differentiation. Deletion of Lrp5 in mice lead to increased expression of Tryptophan Hydroxylase 1, the gut isoform of the enzyme required for serotonin synthesis, thus increasing serum levels of serotonin. Serotonin, in turn, could bind to HTR1B receptors on osteoblasts and stop their proliferation by activating PKA and CREB.

Although different groups have reported controversial results on the existence of an Lrp5-serotonin axis and the action of serotonin in bone remodeling, there is convincing evidence that serotonin modulators such as SSRIs can affect bone turnover. Consequently, the effects of this drug family on periodontal physiology should be thoroughly explored.

Keywords: Serotonin, Alveolar bone loss, Serotonin uptake inhibitors, Periodontitis

Commentary
A recent study by Branco-de-Almeida [1] showed that ligature-induced periodontitis in rats could be ameliorated by Fluoxetine, a selective serotonin re-uptake inhibitor (SSRI), a class of molecules that can increase serotonin levels by inhibiting its clearance inside synapses and are commonly used as antidepressant and as an effective treatment for mood disorders [2]. Carvalho et al., however, showed in this Journal that Venlafaxine, a member of the same drug class, increased bone loss in a rat model of induced periodontitis [3]. These studies raise the question whether and how SSRIs, and therefore serotonin, may affect alveolar bone and the outcome of periodontitis. The issue is of the utmost importance because it could help elucidate poorly known aspects of periodontal pathophysiology in the context of the ongoing debate in bone metabolism, paving the way, if possible, to new therapeutic approaches. As the relation of serotonin and bone is still fiercely debated, the same controversy that has been troubling the bone field seems to be heading for periodontics.

Serotonin
Serotonin, or 5-hydroxytryptamine (5-HT), is a monoamine produced within the central nervous system, mostly in neurons located in the raphe nuclei [4], which send numerous projections to different brain regions, such as the striatum, hippocampus and frontal cortex [5]. Serotonin acts as a neurotransmitter by being released into the synaptic cleft, where it binds to post synaptic receptors. DA transporters (DAT) and 5-HT transporters (5-HTT) collect serotonin from the synaptic cleft and store it in cytoplasmic vesicles in presynaptic neurons, a process commonly referred to as re-uptake, thus regulating the duration of the stimulus. The serotoninergic transmission has a broad spectrum of effects and has been associated to neural
development [6], numerous behavioral and mood disorders [7-11] and to central modulation of pain [12]. Serotonin, however, can also be synthesized by heterochromaffin cells in the gut, where it regulates gastrointestinal function [13], by endothelial cells in the lung [14] and can be found sequestered inside platelet granules [15,16]. As serotonin cannot cross the hematoencephalic barrier, it forms two physically and functionally separated pools, the former inside the central nervous system and the latter in the peripheral body. Although its best known roles are the control of mood, sleep/wake rhythm, peristalsis and mucus secretion, a big but not uncontroversial amount of evidence has been recently reported showing that serotonin may actually exert far reaching effects on bone.

A growing amount of evidence in the literature has also shown that the use of SSRIs is significantly associated to increased risk of fracture [17-23], to increased levels of bone resorption markers [24] and administering SSRIs during pregnancy is associated to shorter length and smaller head circumference in newborns, albeit with unaffected bone quality [25]. Moreover, the frequency of serotonin transporter gene 5-HTTVNTR polymorphism was observed to be higher in osteoporotic patients [26] and 5-HTT polymorphism has been associated to BMD changes after SSRI treatment [21]. Noteworthy, a cross-sectional study by Costa et al. reported an association between aggressive periodontitis and serotonin transporter 5-HTTLPR polymorphism, which has reduced transcriptional efficiency and is associated with lower serotonin re-uptake, in the Brazilian population [27].

The Wnt connection

Our understanding of the role of serotonin has however been considerably increased by studying genetic diseases in human and the effect of alterations in the LRPS5 protein. LRP5 is a membrane protein that is commonly believed to function by dimerizing with transmembrane receptors of the Frizzled family and activating the canonical WNT signaling pathway upon binding to WNT Growth Factors [28-30]. The activation of the canonical pathway requires the recruitment of Disheveled (Dvl) [31,32], which rescues beta catenin from degradation. Beta catenin can be normally found in two pools within the cell, either bound to cadherins in cell-to-cell junctions or in the cytoplasm, where it is sequestered by a destruction complex that targets it for proteosomal degradation [33-36]. Once beta catenin is released, following activation of the receptor complex, it can translocate to the nucleus and bind to a member of the T cell factor/lymphoid enhancer factor (TCF/Lef1) transcription factor family [37,38]. The canonical WNT signaling is required for osteoblast differentiation, bone formation and even bone maintenance through osteoclast inhibition [39-41].

It is well known that loss of function or gain of function mutations in the Lrp5 gene lead respectively to a low or high bone mass phenotype, as clinically observed in Osteoporosis Pseudoglioma Syndrome (OPPG) and the High-bone-mass Syndrome (HBM) [42,43] and the majority of researchers interpret this as the result of Wnt signaling inhibition in osteoblasts. Yadav et al. however showed that deletion of Lrp5 gene in mice lead to increased expression of Tryptophan Hydroxylase 1 (Tph1), the gut isoform that is required for 5-HT synthesis [44]. As a consequence, serum levels of serotonin were increased in Lrp5 knock-out homozygous and heterozygous mice. Their findings are in agreement with numerous clinical observations in OPPG and HBM patients [45-47] and suggest that altered LRP5 functionality may be acting on bone at a distance by regulating circulating levels of serotonin, which would then bind to specific receptors on bone cells, and not directly by controlling Wnt signaling in bone cells, as commonly maintained (Figure 1). Strikingly, ex vivo experiments revealed that 50 mM 5-HT could indeed stop osteoblast proliferation by binding to the HTR1B receptor and activating PKA and CREB transcription factor [44], although previous independent results showed that lower (0.1 mM) doses of serotonin could enhance the proliferation of human primary osteoblasts in vitro [48], suggesting a possible dose-dependent effect. According to Yadav et al. tryptophan-free diet could rescue the bone phenotype in Lrp5−/− mice and, importantly, conditional Lrp5 deletion in the gut increased bone mass [44]. In agreement with these observations, Laporta et al. reported that feeding 5-hydroxy-l-tryptophan to rats from day 13 of pregnancy through day 9 of lactation increased total serum and milk calcium concentrations, osteoclast numbers in bone and bone mRNA levels for resorption markers [49]. Other researchers however have gathered solid evidence supporting the idea that Lrp5 does act locally in bone and does not require serotonin. Niziolek et al. reported an Lrp5 gain-of-function mutation associated with high bone mass in mice without concomitant alterations in circulating serotonin levels [50]. Most strikingly, the same group also showed that global Tph1 deletion did not affect bone mass in their mouse model, in strong contrast to Yadav’s data [51], and Gustafsson et al. reported that administration of 5 mg/kg/day serotonin subcutaneously for 3 months increased Bone Mineral Density and cortical thickness, although reduced trabecular thickness [52].

The pre-clinical observations in rats by Carvalho et al. and Branco-de-Almeida et al. reflect this controversy and do not manage to resolve the conundrum on whether and how 5-HT acts on bone and periodontium. Unfortunately, neither Carvalho nor Branco-de-Almeida considered the role of serotonin in planning their studies
and interpreting their results and as a consequence they did not measure serotonin levels in untreated and treated rats. However, measuring serum serotonin is arguably not an easy task and this could also help explain the discordant results in the literature [53]. Carvalho tested both a low (10 mg/kg/day) and a high (50 mg/kg/day) dose of Venlafaxine per os, whereas only a single but high dose of Fluoxetine (20 mg/kg/day) was used by Branco-de-Almeida, similarly orally administered, so, although it is reasonable to assume that serotonin levels were affected in both studies, it is hard to guess the real extent of it. Moreover, Fluoxetine has been shown to directly reduce osteoblast proliferation and decrease the Osteoprotegerin/Receptor Activator of Nuclear kB Factor Ligand (RANKL) ratio in vitro, thus possibly promoting osteoclastogenesis [48]. Consistently with these results, a 6 month low dose (5 mg/kg/day) treatment with Fluoxetine in rats has been shown to reduce trabecular thickness and increase endocortical bone volume [54]. The dose of Fluoxetine tested by Branco-de-Almeida et al. was on the high end of the doses commonly used to suppress serotonin production in rats [55,56], about twice as high as the maximum recommended human dose (MRHD) [54], and was based on a previous work highlighting its anti-inflammatory properties [57]. It is actually possible that this high dose of Fluoxetine may be activating alternative immunomodulatory or antiinflammatory pathways that are responsible for the authors’ observations and that are overcoming the effects on serotonin metabolism. Indeed Branco-de-Almeida et al. showed that Fluoxetine inhibited IL-1β and COX-2 mRNA and metalloprotease (MMP) 9 activity in their model, and, consistently with these findings, it has been reported that 5 mg/kg i.p Fluoxetine can reduce the expression of MMP 2 and 9 in rat hippocampus [58] and 10 mg/kg i.p. Fluoxetine inhibits the expression of MMP 2, 9 and 12 after spinal cord injury in mouse [59]. A recent study found that SSRIs can differentially control osteoclast and osteoblast viability, apoptosis and activity [60]. Fluoxetine in particular proved to affect preosteoclast viability to a greater extent than other drugs of this class, and this could help explain its effect on periodontitis. Furthermore, it cannot be ruled out that at least part of the positive effects of Fluoxetine on periodontitis could be explained through its action on the central nervous system, where it has been shown that Fluoxetine can increase Tph expression [56,61]. Strikingly, Yadav et al. showed that deletion of the brain stem specific Tryptophan Hydroxylase 2 (Tph2) isoform, lead to a reduction in bone mass in mice, apparently activating an alternative and opposite mechanism to the one mediating the effects of the gut Tph isoform [62]. Both the amount of serotonin and its

**Figure 1 Diagram depicting the two alternative models for Lrp5 action.** According to the former, “central”, serotonin-mediated model (left-hand side), activation of Lrp5 inhibits Tryptophan Hydroxylase 1 (Tph1) and reduces circulating levels of Serotonin, thus decreasing its inhibitory effect on osteoblasts via its HTR1B receptor. According to the “peripheral” serotonin-independent model (right-hand side), Lrp5 acts directly on osteoblasts initiating the canonical Wnt signal cascade and inducing beta catenin translocation to the nucleus and activation of its transcriptional program.
localization appear therefore important for its net effect on bone. Different drugs at different doses might even prevalently act on or have higher affinity for one isoform of the enzyme and the net effect observed on bone could be the result of the systemic suppression of both enzymes.

The controversy is therefore still open and further studies that address this issue in detail are sorely needed.

Perspectives

One of the most exciting aspects of bone mass control by serotonin is the possibility to positively affect bone formation at a distance without acting directly on bone cells, and thus the possibility of a novel therapeutic target to improve the outcome of periodontal disease and alveolar bone regenerative techniques. As of today, a novel compound, LP533401, has been generated and tested in rodents [63,64]. LP533401 can selectively inhibit Tph1 when administered per os, and significantly reduce serotonin levels in blood without passing the hematoencephalic barrier and affect brain’s serotonin concentration. It has been shown that LP533401 can increase bone mass and reduce ovariectomy-induced bone loss in rodents [63,64], even if its active enantiomer, LP923941, was independently proven to be unable to do so [50]. It has not been investigated yet whether LP533401 can affect alveolar bone.

Another equally fascinating aspect of the relation between serotonin and Lrp5 is the possibility, if confirmed, to elucidate some hitherto poorly known regulatory mechanisms in bone physiology. A central dogma of bone biology is that bone tissue is constantly remodeled by teams of dedicated cells that resorb bone, the osteoclasts, and cells that form new tissue, the osteoblasts. These cells act in a tightly coordinated fashion so that bone formation and bone resorption are coupled, because these two processes occur simultaneously and affect each other. It has been shown that osteoclasts can mobilize sequenced osteogenic factors such as TGFβ1 from the mineralized matrix during resorption, stimulating osteoblasts and presumably contributing to sustain bone formation in the osteoclastic lacuna [65]. It is also known that cells of the osteoblastic lineage can produce downstream, such as possibly serotonin, could indeed unravel the secret to uncoupling bone formation and resorption, with huge therapeutical benefits.

Although the available results in rodents are just preliminary, they are undoubtedly captivating and are an open invitation to further investigate the role of serotonin not only in bone physiology but also on periodontal physiology and pathology.

Competing interests

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Authors’ contributions

GC conceived the manuscript and drafted it. PG helped GC drafting the manuscript and acted as consultant on bone physiology. MG acted as consultant on serotonin physiology and SSRI antidepressants. All authors read and approved the final manuscript.

Authors’ information

GC, DDS, PhD is mainly interested in alveolar bone regeneration and new approaches to promote the healing of bone defects. PG, MD, PhD is a bone specialist who focuses on bone loss and osteoporosis. MG, MD, DDS, PhD was trained as a neurologist and a periodontologist. His main research focus is oral neuropathophysiology, TMD and sleep disorders.

Author details

1. Dep. Biomedicine, Biotechnology and Translational Sciences, University of Parma, Via Gramsci 14, Parma 43126, Italy. 2. Dep. Clinical and Experimental Medicine, University of Parma, Parma, Italy.

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