Human uterine leiomyomas (ULs) being the commonest neoplasms affecting women of reproductive age affect African American (AA) much more as compared to Caucasian (CC) women. One of the possible reasons thought is in view of higher incidence of hypovitaminosis D in this group. We have been trying to look for a permanent medical agent that can be used long-term for fibroid treatment instead of the earlier selective progesterone receptor modulators (SPRM)'s of which ulipristal acetate has been recommended by the FDA but in view of its side effects like progesterone associated endometrial changes (PAECs), it can't be used for long-term. Thus, in an endeavor to find a medical therapy that can be used for long term without any adverse side effects Vitamin D is getting pursued with its effect on Wnt/beta-catenin pathway by which it can affect the MED12 status, a possible cause of etiopathogenesis of UL. Human uterine leiomyomas (ULs) affect roughly 70% of women of reproductive age [1], being the most common benign estrogen-dependent tumors that develop during the reproductive period [2,3]. It is most prevalent in African American (AA) women (73%), in comparison with Caucasian (CC) women (45%) [4]. Though they are benign, nonetheless these tumors cause significant morbidity, being the main cause for hysterectomy and also for both gynecologic dysfunctions associated with reproductive dysfunction that varies from heavy menses along with pelvic pain to infertility, recurrent miscarriage and preterm labour. The etiopathogenesis of ULs is mostly not clear, yet MED12 mutations are the most common driving mutations accounting for roughly 70% of ULs [5,6].

The canonical Wnt/β-catenin pathway is a well characterized cell signaling pathway, connected to the formation of many kinds of tumour. The activation occurs downstream of the interaction of a Wnt ligand with the Frizzle/LRP co-receptor complex, that prevents the degradation of β-catenin targeted genes in human myometrial cells [7]. Earlier Al-Hendy et al. [7] showed that β-catenin signaling was activated in the myometrium of ULs with MED12 mutation in comparison of the normal adjacent myometrium. The MED 12 G>A mutations cause a raised expression of β-catenin targeted genes in human myometrium cells [7]. A paracrine role for Wnt/β-catenin signaling has been proposed by Ono et al [8] that helps differentiated myometrial cells or UL cells to send mitogenic signals to the nearby myometrial stem cells in response to estrogen (E)and progesterone(P), that causes growth of ULs. A woman has a threefold greater incidence rate and relative risk of ULs than Caucasians (CC)women. Basis of this risk differentiation is not clear, some current studies point to hypovitaminosis D as the main culprit, since AA women have a ten-times >risk of Vitamin deficiency as compared with CC women [5]. With these findings with the more recent observations that vitamin D3 and its potent nonhypercalcemic analog paricalcitol effectively depressed UL growth in animal models [5], implicates that analogs of Vitamin D and Vitamin D might be promising to prevent or for early therapy. Further Al-Handy et al. [7] demonstrated that silencing of Vitamin D receptor (VDR) in normal myometrial cells induces expression of Wnt/β-catenin and stimulates a fibrotic process, which involves raised cell proliferation along with extracellular matrix production [9].
Recently Corachan et al. [10] evaluated the differential expression of proliferation and apoptosis-related markers like PCNA, BCL2 and BAX along with WISP1 (Wnt inducible signaling pathway protein) in various matched myometrial and UL tissues. Initially they showed that the upregulation of PCNA and WISP1 was seen in ULs that is similar to the previous observations. But they did not find impairment in apoptosis in ULs they saw, probably patient variabilities might be a contributing factor. Nevertheless, on evaluation of these markers in matched patient tissues is definitely more reliably. They then examined the effect of Vitamin D on induction of isolated primary cell growth arrest along with decrease in PCNA protein expression. With the use of primary cells to measure Vitamin D effect is a point that adds strength in contrast to immortalized cell lines. Furthermore, Wnt signaling along with apoptosis specific polymerase chain reaction arrays have been utilized for determining the Vitamin D response on gene expression which are involved in crucial pathways. More in depth studies might be carried out from observing these studies in future.

Though the efficiency of Vitamin D therapy is correlated with patient disparity along with levels of VDR, the ethnicity of the collected patient samples was not described in this article. If the observations related to the levels of VDR in ULs are present in this study one will obtain more detailed information regarding the efficiency of treatment with vitamin D3 in ULs. Since this correlation has been ignored, this relation might explain why apoptotic effect of Vitamin D treatment on isolated primary cells in this study is not consistent with the earlier findings that showed that Vitamin D therapy causes apoptosis in UL cells [5], since VDR is a nuclear receptor which responds to its ligand (i.e. Vitamin D), as demonstrated earlier [5,9]. Hence need for dividing subgroup into the samples tested as per VDR expression might demonstrate marked effect following Vitamin D therapy. Likewise testing the MED12 mutation status in the tissues might give helpful knowledge for understanding the Wnt polymerase chain reaction arrays data once Vitamin D therapy has been given, in view of MED12 mutation has been demonstrated to modulate Wnt/β-catenin signaling. Hence pooling all the samples may dilute any effect and lead to minimal changes of gene expression secondary to Vitamin D therapy. Further measuring the protein levels of particular markers might have greater impact. Nuclear translocation of β-catenin instead of expression levels have been demonstrated to be responsible for etiopathogenesis of particular cancers. Hence measurement of expression of nuclear β-catenin expression in ULs might indicate the activation of this particular pathway more adequately. Further changes in β-catenin’s expression i.e., decreased levels after testing efficacy of treatments like Vitamin D as anti-UL therapy of the modalities available. Moreover, since ULs are derived from monodonal stem cells, genome wide analysis of Vitamin D’s effect is needed. Earlier we reviewed the different approaches for medical management of fibroids [11], instead of the surgical or radiologic management which are not desired by the patient specially if she wants preservation of fertility in future. Hormonal therapies are designed to blunt the growth stimulatory actions of E and P on UL growth like selective progesterone receptor modulators (SPRM) which are approved for a short time in view of complications like progesterone associated endometrial changes (PAEC) on long term use. Ciebiera et al. [12] reviewed how Vitamin D might be obtained and its metabolic pathways (Figure 1). Further they showed the influence of Vitamin D on UL (Figure 2). Further they compared effects of Vitamin D, Gn RH analogs and SPRM’S (Figure 3). As per Ali et al there is a potential synergism between Vitamin D and ulipristal acetate [13,14]. Further Vitamin D also has an effect on extracellular matrix implicated in UL pathogenesis [15]. Thus, for long term therapies that are targeted for effective UL size decrease and ameliorate the common symptoms are actively needed for addressing this. These would not be affecting future fertility and would be safe and cost effective in view of natural origin of Vitamin D and its availability, hence it might appear the safest, cheaper option.

**Figure 1:** Courtesy [12]. Vitamin D metabolism and schematic pathways. Diet, supplements and sunlight are the major source of vitamin D in humans. Vitamin D is synthetized in skin from 7-cholesterol. In further steps liver converts it to 25(OH)D and then kidney to 1,25(OH)D. Optimal vitamin D serum levels were described as 25(OH)D of 40-60ng/mL.
Figure 2: Courtesy [12]. Vitamin D influence on uterine fibroid (UF) pathophysiological pathways. Cyclin-dependent kinase 1 (CDK1), proliferating cell nuclear antigen (PCNA), catechol-O methyltransferase (COMT), Bcl-2 protein, proliferation marker protein Ki-67 (MKI-67), extracellular matrix (ECM), transforming growth factor beta 3 (TGF-β3), uterine fibroid (UF).

Figure 3: Courtesy [12]. Vitamin D, GnRH analogs and SPRMs. According to Ali et al. there is a potential synergism between vitamin D and ulipristal acetate [140]. New data about between vitamin D, vitamin D analogs and other mostly used drugs in UF therapy is necessary to find other synergisms. Gonadotropin releasing hormone (GnRH), selective progesterone receptor modulator (SPRM).

Conclusion

UL’s being one of the commonest neoplasms in women of reproductive age require treatment. Earlier TAH was considered the only treatment or operative myomectomy/laparoscopic one was the only answer. After decade of trying for development of medical management still we have not managed to develop an effective treatment with SPRM, the commonest used of which ulipristal acetate has been approved but still flaw is it can’t be used for long term in view of side effects like PAEC. One of the major proposed causes of development of UL’s is MED12 mutations and Vitamin D has been shown to act via Wnt/β-catenin pathway to affect these uterine leiomyomas. Further studies are going on to ensure that this is the permanent medical treatment for UL’s we have been looking for.

References


