Purpose of study: Understanding that the inhibition or mutation in S6K and in hydrophonic acids (Ser, Tyr, Pro...), in BTK and in PLCγ1 will lead to inhibition in PLCγ2 and in Thromboxane-A2 then will be the main reasons for chronic lymphocytic leukemia “CLL” disease, where proper S6K/BTK and PLCγ2 are main regulations for thromboxane-A synthesis and necessary for B-cells maturations and T-cells modulations.

Also, it’s important to understand main factors that cause and link the Osteoarthritis “OA” with diabetes which are the deficiency in Ser (hydrophobic) amino acids and the mutated S6K productions lead to deficiency or inhibition in Ser phosphorylation signaling which normally is the basis of Ser/Thr phosphorylation signalling which necessarily for Akt, for S6K1 synthesis and necessary for RORs and IFNs synthesis, and also necessary for proper PLCγ2 productions, where S6K is the main regulator for ATPase, for ribosomes, for OPA1 repair, and for proper PLCγ2 synthesis, that I have to note that the percentage of the shortages ratio of amino acids or in the increasing in positive linkages are the main ratio that can define the degree and type of specific disease which can differ from other diseases or can linked with the same Syndromes of other diseases. That also the shortage ratio between the beta Cytokines productions and the ratio of sudden high inflammations productions “and the type of its inflammatory molecules” have to be calculated and considered related to the patients ages (whether child, youth or old ages) and the duration of the chronic disease disease, that some can be confused to differentiate between auto-immune disease and regular disease problems diagnosis.

That, There was a case of a child with 9-year-old who had a suspicion of loose of bone maturation and growth and has a sudden infection in the right lung and a lack of breathing with pain. It was found that there was a pulmonary abscesses in right lung and there was a development with the appearance of an air bag or “inflammatory fluid bag” surrounding respiratory cells in right side. The occurrence of sudden inflammations molecules and their growth was rapid enough faster than IFNs productions and faster than PLCγ2 productions due to the age of the child, “Note some her regular treating doctors diagnosed her medical conditions as a type Autoimmune disease and she has weakened immunity due to sudden fast infection related to her young age”.

Highlights:
_ Increasing in PLCγ1 with Deficiency in Ser, deficiency in proper S6K, and decreasing in synthease activity with inhibition in PLCγ2 will reflect decreasing in anti-inflammatory processes, reflect Osteoarthritis and diabetes syndromes, and also inhibition in PLCγ2 will inhibit TXA2 synthesis and can be the main reason for CLL diseases.

_proper healthy PLCγ2 are so necessary for increasing re-medulate immune efficiencies, and for re-modulate IgM and IgD antigen and T-cells functions, and also proper healthy PLCγ2 productions (which depend on PLCγ1 and BTK Biosynthesis) are so imp for recover osteoporosis and both Osteoarthritis and diabetes.

_inhibition in PLCγ2 Bio-Synthesis can reflect decreasing or inhibition in Thromboxane-A1 percentages and can lead to CLL diseases, Where, CLL characterized by inhibition in BTK which regulate PLCγ2 synthesis, inhibition in main antigen synthesis, and then inhibition in the proper normal Thromboxane-A synthesis (which regulated mainly by PLCγ1 and by BTK and then linked to IFNs productions and regulated by PLCγ2 proper productions).
Chronic lymphocytic leukemia (CLL) observed during treatment with B-cell receptor inhibitors pathway including inhibitor of Bruton’s tyrosine kinase-PLCγ2, where, CLL can be strongly linked to Osteoporosis “OA” and Linked to both Osteoarthritis and diabetes too.

**Keywords:**
- Phospholipase C-1 “PLCγ1”
- Phospholipase C-2 “PLCγ2” “necessary for anti-inflammatory steps”
- Osteoarthritis OA tissue cells
- Osteoporosis tissue cells
- Osteoelast processes Osteoblast processes
- Ser/Thr phosphorylation signal
- Deficiency in PS/-Thymine-kinases reflect mutated S6K, deficiency in PLCγ2, deficiency in B-cells and T-cells modulation, and deficiency in OA1 repair
- S6K, estrogen, androgone, JAK state signaling
- Diabetes pathogenic tissue cells
- Tyrosine phosphatase

**Abstract:**
Proper S6K /BTK and PLCγ2 synthesis (which regulated by Ser phosphorylation pathway) are main regulations for thromboxane-A “TXA2” synthesis, and necessary for B-cell maturations and T-cells modulations and functions.

The main reasons for causing Osteoarthritis “OA” and diabetes diseases (that are linked together) are the deficiency of Ser amino acids and decreasing of Ser phosphorylation signalling pathway which necessary for proper S6K productions, where normally the Ser phosphorylation signalling pathway is the basis of Ser /Thr phosphorylation signalling and is necessary for proper Akt, S6K1 synthesis and necessary for RORS and IFNs synthesis and also necessary for running proper BTK for PLCγ2 productions, where S6K is main regulator for ATPase and for proper PLCγ1 and for PLCγ2 synthesis which necessary for bone growth and for modulating immune efficiency.

Osteoarthritis “OA” is characterized by a sharp expression in Gamma-Phospholipase C-1 “PLCγ1”, with decreasing “or inhibition” in PLCγ2 “PLC beta” productions due to inhibition or mutation in S6K and then in BTK.

The increasing in PLCγ1 with Deficiency in Ser amino acids will lead to deficiency in Ser phosphorylation signalling, and decreasing in synthase activity that will reflect down regulations in BTK pathways and lead to inhibition in PLCγ2 productions which will reflect diabetes (inhibition in Estrogen with the production of Androgen instead of estrogen) and can reflect Osteoarthritis “OA” prognosis depend on the percentage of Deficiency or inhibition in basic amino acids and its basic necessary signaling pathways.

The proper S6K are so necessary for reactivating both PLCγ1&2 , where phospholipase Cy2 (PLCγ2) is activated from a variety of cell surface receptors such as Syk “S6K”.

The B-cells are promoted by the function and activities of both PLCγ1&2, but the deficiency in Ser amino acids will reflect decreasing in Ser phosphorylation pathways and then decreasing in Estrogen synthesis, with increasing in Androgen synthesis which lead to decreasing in PLCs isoforms production and lead to pathogenic diabetes problem. So T2DM is strongly connected with OA diseases are linked together that deficiency in Ser amino acids and their phosphorylation can cause both diseases and any early step from any of those two or more similar diseases can lead to the other.

Pathogenic type 2 diabetes associated with progressive beta-cell impairment due to the mutations in the production of S6K1 and inhibition in the PLCγ2 which due to inhibition or decreasing in Ser phosphorylation signaling during mTOR Ser/Thr phosphorylation pathways that can reflect Inhibition in The releasing of PS/T-Thymine Kinases and PS/T-Cytosine kinases chains (mTORC1) which are depending on availability of hydrophobic amino acids synthesis including Ser and Tyr which can be modified by synthetase enzymes for creating active gamma-subunits (upon synthetase effects) that can be modified by synthase effect for Beta-subunit synthesis “PLCγ2” then will be modified by phospholipase effects for alpha subunits productions.

The releasing of PS/T-Thymine -Kinase and PS/T-Cytosine -kinase chains (mTORC1) from the phosphorylations of Ser amino acids is so necessary steps for normal proper S6K productions, which necessary for IFN-Gamma and for PLCγ1 productions, and therefore necessary for normal PLCγ2 synthesis upon “BTK activity” which is necessary for B-cell maturations, for T-cells modulations, for modulating anti-inflammatory steps and procedures, for thromboxane-A synthesis, and for bone growth and modulation.

Inhibition in PS/T-Thymine -Kinase and PS/T-Cytosine -kinase chains (mTORC1) from the phosphorylations of Ser amino acids is so necessary steps for normal proper S6K productions, which necessary for IFN-Gamma and for PLCγ1 productions, and therefore necessary for normal PLCγ2 synthesis upon “BTK activity” which is necessary for B-cell maturations, for T-cells modulations, for modulating anti-inflammatory steps and procedures, for thromboxane-A synthesis, and for bone growth and modulation.

It’s imp to note that Tyrosine phosphatase PTPs are important regulator of chondrogenic patterning and are critical regulators of tyrosine phosphorylation that it’s activity depends on Tyr, Ser synthesis (hydrophobic acids) and on JAK state signaling activities.

And so, the proline-rich tyrosine kinases regulate proper PLCs isoforms which compete for binding site at the very C terminus of fibroblast growth factor for osteorogenitor embryonic development , and bone formations.

Synthetase is the main regulator for PLCγ1 activities followed by synthase effects which is the main regulator for beta-subunits “PLCγ2” productions which is able to “upregulate phospholipase abtivity” for alpha subunits (PLC-alpha) productions for reactivating fibroblast growth factor receptor (FGFR2), for reactivating both IgM and IgD and for TLR4 productions for osteoblast processes.

Where, PLCγ1 competes for a binding site at the very C terminus of FGFR2 for embryonic development and bones growth, where, PLCs isoforms are involved in multiple stages in TLR4, interferon, and in anti-inflammatory steps.
And also, PLCγ1 recruit to CSF-1 is following imp stages for producing PLCγ2 which is necessary for activating anti-inflammatory where, IFN-γ activates PLC-γ2 via an upstream of tyrosine kinase.

PLCγ1 recruited to CSF-1 for two pathways activities 1st / re-activating IFNs productions which regulate MHC class1 and class two for modulating cell-surface protein activities, 2nd / activating PLCγ2 for modulating T-cells, where PLCγ1 involved in the production of TRIM22 for mediating antiviral activities and anti-inflammatory processes through reactivating IFNs productions for PLCγ2 synthesis.

PLCγ2 are so imp in anti-inflammatory processes and can be considered as having the main roles for thromboxane-A synthesis.

Inhibitions or mutations in S6K, in BTK and then in PLCγ2 productions will cause an inherent or inhibition in CXCL12 then followed by inherent or inhibition in CXCR4 then reflect inherent or inhibition in the regulation of B-cell growth through mutations in IgM and in IgD.

Proline amino acids is necessary for reactivate OPA1 anabolic oxidations (started by synthetase, then synthase, then phospholipase for producing gamma “PLCγ1”, then beta “PLCγ2”, and then alpha “PLC-α” subunits respectively) for cartilage synthesis which promote PLCγ2 synthesis necessary for bone growth including antigen and thromboxane-A synthesis.

Introduction:

Osteoarthritis is characterized by a sharp expression in Gamma-Phospholipase C-1 “PLCγ1”, with decreasing in PLCγ2 “PLC beta” which improved by phospholipase oxidations for producing PLC alpha which necessary for proliferations and calcium entry “, where PLCγ1 was highly expressed in human OA chondrocytes [1] ) which is implicated processes including mitogenesis and calcium entry.

Phospholipase C isoforms (PLCs) are essential mediators for cellular signaling and for cellular metabolism. PLCs regulates multiple cellular processes including proliferations and biological bones growth by generating bioactive molecules such as inositol-1,4,5-triphosphate (IP3) and diacylglycerol.

That, PLCγ1 basis of inhibition-driven autophagy of IL-1β-treated chondrocyte confers cartilage protection against osteoarthritis. [2] the only presence PLCγ1 has the roles of analyzing biological molecules “Osteoclast” through expressing its own specified functions, while PLCγ2 has the role of functioning PLCγ1 for running anti-inflammatory processes and for promoting proliferation through activating phospholipase for activating PLC alpha for running proliferation.

So slightly inhibition or decreasing in PLCγ1 will decrease osteoclast and also re-functioning PLCγ1 for reactivating the expression of PLCγ2 “which reduce the analyzing function of PLCγ1”, then will give the priority for PLC-beta “PLCγ2” for production by beta-oxidations for activating anti-inflammatory processes, and for promoting PLC-alpha production which necessary for proliferations functions that will activate osteoblast processes, bone growth, and cells proliferation “modulations”. The availability of Proline amino acids are necessary for activating and accelerating OPA1 oxidative processes which activate cartilages synthesis through PLCγ2 synthesis, and the necessary of hydrophobic amino acids availability and proper synthesis in vivo is important for gamma subunits synthesis regulated by synthetase enzymes “eg : Tyr, Leu, Pro, Gly, Ser, ... etc”. Proline with hydroponic acids can activate and accelerate proper OPA1 oxidative processes which promote and activate necessary anabolic cycles by activating BTK which regulate PLCγ2 synthesis and bone growth and for modulating immune effectiveness.

The Deficiency in the conversion of glutarate to glutamate and decreasing in proline biosynthesis strongly affect on cartilage synthesis due to decreasing in the activation of mitochondrial OPA1 oxidative processes. [3] Also, deficiency in the mitochondrial OPA1 membrane repairs process of can reflect deficiency in the proper S6K productions (which necessary for ATP and GTPase synthesis where GTPase is strong regulator for OPA1 repair ) that will lead to deficiency in OPA1 mitochondrial function that will lead to decreasing in PLCs synthesis (decreasing in PLCγ2) then in SIRPα1, and in TLR4 biosynthesis, that can reflect increasing in catabolic analyzing processes (due to increasing in PSTG-kinases and PSTA-kinases “for a limit” with decreasing in PS / T Thymine kinases and in PS/T-Cytosine kinases productions which will lead to diabetes, where in diabetes the synthetase activities can be increased that can analyze phospholipids, foreign molecules and Enterstatium fluid lead to decreasing in the synthesis of anti-inflammatory tools eg PLCγ2. Proper S6K1 synthesis can promote ATPase and GTPase productions through OPA1 repair for activating RORs pathways, for activating BTK pathways and for PLCγ2 productions, where all are depending on the four types of kinases production from Ser Thr phosphorylation pathways which activate proper PLCs productions, proper MIHs synthesis and proper bone growth synthesis with beta-cells proper maturations and T-cells modulations.

Method and results:

Proper S6K / BTK are regulating PLCγ2 synthesis and are regulating proper thromboxane-A synthesis, B-cell maturations and T-cells modulations.

Where, it’s so important to Understand main factors that cause Osteoarthritis “OA” and diabetes which are the deficiency of Ser amino acids which lead to mutated S6K production due to deficiency or inhibition in Ser phosphorylation signaling which normally is the basis of Ser /Thr phosphorylation signalling that are necessary for proper Akt, S6K1 synthesis and necessary for proper MHCs synthesis and proper bone growth synthesis with beta-cells proper maturations and T-cells modulations.

Proper S6K productions through availability of Ser are main regulator for ATPase synthesis and GTPase which necessary for OPA1 repair, and for BTK pathways and proper PLCγ1 synthesis which regulate PLCγ2 synthesis for necessary bone growth and cartilage synthesis.

Osteoarthritis “OA” is characterized by a sharp expression in Gamma-Phospholipase C-1 “PLCγ1”, with decreasing “or inhibition” in BTK which lead to decreasing in PLCγ2 “PLC beta” that will reflect decreasing in beta-cells maturation and functions and in T-cells modulations.

The increasing in PLCγ1 with Deficiency in Ser will lead to mutated S6K production, and decreasing in proper synthase activity and decreasing in BTK processes that will lead to inhibition in PLCγ2 synthesis that will reflect Inhibition in Estrogen synthesis and increasing Androgyne synthesis that will give the Symptoms of diabetes and Osteoarthritis "OA" diseases.

we’ll discuss why both diseases are connected and their causes depend mainly on availability of Ser and hydroponic amino acids, and depend on the proper S6K synthesis then on the Tyr and other hydroponic acids synthesis and their phosphorylation.
signaling pathway and JAK for the Synthesis of their Receptors. Deficiency in the proper S6K, in Ser and in Tyrosine synthesis “regulated by synthetase” will lead to increasing in PLCγ1 with decreasing in PLCγ2 synthesis (which are Regulated by availability of PS /-Cytosine k and PS /-Thymine Kinases) will lead to androgen synthesis instead of estrogen “diabetes” and Osteoarthritis “OA” diseases:

PLCγ1 is a protein molecules that its activity depending on Tyr phosphatase and gamma common receptors synthesis which regulated by JAK STAT signaling, and also regulated by synthetase enzyme where synthetase is the main second enzyme in OPA1 chains after COX enzyme (followed by synthase and phospholipase respectively ) and necessary for hydroponic acids synthesis for gamma active subunits synthesis (or extraction), that synthetase enzymes is so necessary for creating signals transmission which can reactivate mTOR Ser/Thr signaling pathway and for re-producing the active gamma-subunits which upon JAK signaling will produce their active receptors necessary for activating gamma subunits “PLCγ1” beta-subunits “PLCγ2” synthesis which upon phospholipase will produce alpha subunits “PLC-alpha” for activating proliferations, and bones growth. The PLCγ1/PLCγ2 double-deficient B cell progenitors have reduced expression of genes related to B cell lineage, IL-7 signaling, and cell cycle. [4] That the activities of both PLCγ1&2 are linked to each other and are so necessary for re-activating B-cells maturation, where, PLCγ2 regulate the productions antigen-specific immunoglobulin necessary IgM and IgD synthesis necessary for anti-inflammatory processes, and necessary for T-cells modulations, therefore the deficiency or mutations in PLCγ1&2 will lead to decreasing in or lead to Malignant transformation in B cells that can cause mutations or inhibition in IgM and in IgD synthesis and will lead to inhibition in mutations in TXA2 synthesis too that will lead to a cancer problem as chronic lymphocytic leukemia (CLL) and can cause several other pathogenic problems as diabetes and OA diseases.

B-cells are promoted by the productions of both PLCγ1 which upon BTK will regulate PLCγ2 synthesis, where PLCγ1 synthesis mainly depends on mTOR Ser /Thr phosphorylations signalling pathways (mTS/Tp ) and depend on proper S6K synthesis “that deficiency in Ser amino acids will reflect decreasing in the productions of the two types kinases PSTT-K and PSTC-k that will lead to mutations in S6K synthesis and lead to decreasing in Estrogen synthesis with increasing in Androgen synthesis which lead to pathogenic diabetes diseases.[5]

Proper S6K synthesis is depending on availability of Ser amino acids and on the production of the two kinases PSTT-K and PSTCk that are so necessary for reactivating ribosomal ATPase which is necessary for repairing the mitochondrial OPA1 membran (through regulating GTPase productions), where proper OPA1 is necessary for activates and regulating proper PLCγ1 productions and for “PLCγ2” synthesis upon synthase effect for B-cell receptor synthesis for B-cells maturation, and for anti-inflammatory, then followed by creating PLC-alpha synthesis upon phospholipase functions for promoting proliferations and bone growth through SIRPα and TLR4 productions.

In case of deficiency in mTOR Ser/Thr phosphorylations signalling due to deficiency in Ser phosphorylation will produce non proper mutated S6K “missing Ser hydrophobic amino acids” that will lead to diabetes pathogenic problems, and will lead to inhibition in PLCγ2 or will lead to mutated PLCγ2 in some cases depending on the percentage of Deficiency of necessary hydroponic (Tyr, leu, Pro,... etc) that will lead to inhibition in Estrogen which is the substrate for RORs pathways and will lead to increasing in Androgyne instead of Estrogen that will inhibit PLCγ2 synthesis and will lead to diabetes, and OA diseases and can lead to cancer pathogenesis in the inhibition or mutation in TXA2 synthesis.

Pathogenic type 2 diabetes associated with progressive beta-cell impairment due to the mutations in the production of normal Estrogen (that Androgyne is formed instead of Estrogen) due to missing of Ser phosphorylation signaling during mTOR Ser/Thr phosphorylation pathways due to Inhibition or decreasing in the releasing PS/T-Thymine -Kinases and PS/T-Cytosine -kinases (mTORC1) where those two kinases synthesis depending on the availability of Ser amino acids and depend on Ser phosphorylation pathway, (where synthetase enzymes regulate hydrophobic amino acids synthesis 8n vivo) where through gamma oxidations by synthetase will promote gamma-subunits that will be modified by synthase effect to produce active Beta-subsunits synthesis which are necessary for “anti-inflammations and for promote alpha subunits synthesis upon phospholipase effects “alpha-oxidations” which necessary for proliferation respectively.

The releasing of PS/T-Thymine -Kinase and PS/T-Cytosine -kinase chains or (mTORC1) from specifically the phospholylations of Ser signalling pathway is so necessary for the mechanism of normal and proper S6K productions which necessary for ATPase synthesis, for IFN-Gamma productions, and for activating BTK which necessary for promoting PLCγ2 productions which is necessary for B-cell productions and functions , for T-cells modulations, for modulating anti-inflammatory steps and processes , for thromboxane-A synthesis, and for bone growth and maturation.[6*]

The inhibition in PS/T-Thymine-Kinase and PS/T-Cytosine -kinase (mTORC1) productions will be the main reason for the inhibition of the beta subunits productions “PLCγ2” that can be the reason of decreasing in the hyperpolarization and then electrical activity will lead to decreasing in the abolition of Ca2+ which will lead to decreasing in blood pressure and Ca2+ precipitations in blood vessels.

Also, the deficiency in Tyrosine amino acids will prevent the production of tyrosine phosphatase which needed for the synthesis of phospholipase C 1&2 that promote cellular proliferation, and the reduction and deficiency in Tyr amino acids “hydrophobic acids” will reduce or inhibit Drutons tyrosine kinases “DTK”. Now it is important to consider that proper S6K is the main regulator for PLCs isoforms synthesis which depend on S6K productions, and it has been reported that the phospholipase C γ2 (PLCγ2) is activated from a variety of cell surface receptors such as Syk “S6K”, and BTK which phosphorylate and activate PLCγ2 [6].

Proper S6K1 synthesis is the basis for ATPase, and GTPase synthesis and also is the basis for ribosome repair where, GTPase is necessary for G-protein synthesis, for OPA1 membrane repair, and for ribosomal repairs that always necessary for regulating cellular growth and anti-inflammatory Processes. As the GTPase is a regulator tool for BH4 and NO 3 productions, and it has been reported that the phospholipase Cγ2 (PLCγ2) is activated from a variety of cell surface receptors such as Syk “S6K”, and BTK which phosphorylate and activate PLCγ2 [6].
Tyrosine, Ser, and proline are necessary hydrophobic acids such as Tyrosine, Ser, proline are necessary growth. Productions, where PLCγ2 is also regulated by BTK for proper creating necessary receptors for both PLCγ1 and then PLCγ2 OPA1 synthetase and then activated by JAK STAT signaling for mTOR Ser /Thr signaling pathway that will not produce normal S6K “due to deficiency in Ser and some other necessary amino acids (mainly Ser, Tyr, Leu, Pro a.a.) then will lead to decreasing “or mutation” in the S6K productions, that will lead to Androgen production instead of Estrogen where Estrogen characterized by presence of Ser in their molecules “and is the substrates for ROR anabolic signaling pathways”, that will lead to high ATPase productions (due to availability of purines with decreasing in pyrimidine synthesis) with deficiency estrogen synthesis that later will promote the IFN gamma, IFN-beta, and alpha that can lead to increasing in “catabolic processes” with decreasing in the ROR pathways “anabolic process” and decreasing in proper PLCγ2 productions that will reflect Ca⁺⁺ precipitations and arterial hypertension.

Where, it has been reported that insulin activates the K-ATP channels of pancreatic β-cells and islets, resulting in membrane hyperpolarization, and the abolition of [Ca²⁺]i oscillations [8]. And, the low abolition of [Ca²⁺]i oscillations in the case of T2DM indicates decreasing or inhibition in pyrimidine synthesis “regulated by synthetase”, decreasing in synthase functions, and decreasing in PLCγ2 synthesis “that has the role of modulating inositol 1,4,5-trisphosphate-mediated calcium oscillations for bone growth”. Also, decreasing in membrane hyperpolarization can give reflection of decreasing in OPA1 synthase oxidations which reflect decreasing in membrane hyperpolarization and decreasing in PLCγ2 synthesis.

(PLCγ1) can be reactivated by platelet-derived growth factor “GF” receptors, insulin-like GF 1 receptor (which reflect deficiency in proper cells and bones growth), but in brief PLCγ1 productions can produced and re-functioned by several active growth factor (GF) receptors through feedback and by firstly reactivating synthetase followed by synthase then phospholipase which promote growth factor activities as epidermal GF receptor [EGFR], and platelet-derived GF receptor, where due to activating GFs processes it will be responsible for increasing hyperpolarization and functioning CA throughout the synthesis of PLCs that will responsible for running the pathway of bone growth and cellular biosynthesis processes.

The main PLCγ1 proper activities is regulated firstly by main ribosomes and by proper S6K productions from mTOR Ser /Thr phosphorylation pathways followed by JAK STAT signaling for producing the Tyr-phosphatase, gamma common receptors, and other necessary helical proteins receptors which adopt and activate PLCγ1& 2 synthesis and activities for anti-inflammatory, for B-cells maturation, for T-cells modulation, and for bone growth and proper cellular proliferation.

PLCγ1 is a necessary Protein regulated firstly by chromosomes, then by ribosomes activities and by S6K which produced from mTOR Ser /Thr signaling pathway that regulated firstly by OPA1 synthetase and then activated by JAK STAT signaling for creating necessary receptors for both PLCγ1 and then PLCγ2 productions, where PLCγ2 is also regulated by BTK for proper PLCs isoforms productions for cellular proliferation and bones growth. Hydrophobic acids such as Tyrosine, Ser, proline are necessary for facilitate the cellular and B-cells maturation and survival that protect proliferation processes of bones development (also can activate tumor growth in case of synthase dysfunction when lose or deprived of some necessary amino acids) through facilitating OPA1 oxidative functions (that proline is necessary for OPA1 enzymes activities which activate their function and bone cartilage growth ) and activate BTK pathways which necessary for FGFR2 gene expression for bones developments. Where, Tyrosine amino acids increase alertness and bone development through activating tyrosine kinases, that Tyrosine phosphatases which are potential therapeutic targets for fighting bone disorders [9].

Protein tyrosine phosphatase (PTP) gamma (carry→ve charge regulated firstly by synthetase gamma-oxidations) has been proposed to be an important regulator of chondrogenic patterning, where PTPs are critical regulators of tyrosine phosphorylation at multiple stages of bone development and metabolism [10]. And, proline-rich tyrosine kinases regulate osteoprogenitor cells and bone formations, [11] so Tyrosine and Proline (where their synthesis firstly regulated by synthetase in vivo) are regulated by PIPs and are critical regulators for multiple stages in bone development started by cartilage synthesis.

Tyrosine, Ser and proline are essential hydrophobic acids that produced in vivo upon the effects of synthetase enzymes on nutrients-mTOR , and on inflammations molecules for running pyrimidine synthesis for creating and improving (modulating) active Gamma-subunits for PLCγ1 synthesis which modulated and regulate the beta subunits “PLCG2” (upon BTK regulation) which necessary for increasing and modulating anti-inflammatory efficiency , then the PLCγ2 will be modulated for producing alpha “PLC-alpha” active subunits productions which necessary for proliferation, B-cells maturations, and bone growth. Gamma-subunits firstly moderated by JAK STAT signaling for producing their own active gamma subunits receptors (as Gamma-common and other helical proteins) which can be promoted by IFN gamma too for re-activating PLCγ1, PD-1, MHC-class-1 and class two, (where PLCγ2 promote antigens IgM and IgD), then MHC class two promote the SIRPα1, TLR4, and PD-L1 productions necessary for bone growth, cells developments and T-cells modulations.

PLCγ1 competes for binding site at very C terminus of FGFR2 for embryonic development and bones growth, where, PLC isoforms are involved in multiple stages in TLR4, and in interferons production: PLCγ1 competes for a binding site at C terminus of fibroblast growth factor receptor (FGFR2) (which plays an important role in bone growth, particularly during “embryonic development” and is sufficient to upregulate phospholipase activity [12]. That, S6K and synthetase regulate PLCγ1 production followed by BTK and synthase effects for beta-subunits (“PLCγ2”) productions which stimulate phospholipase “ablivity” for up regulating phospholipase activity for active alpha subunits (PLC-alpha) productions which can reactivate the production of fibroblast growth factor and their receptors (FGFR2) for full proliferations cycles, bone growth, cells maturation and T-cells modulations. Their are strong relationships between PLCγ1&2 bio-activities and productions of the MHC class 1 and two which promote SIRPα1, TLR4, and PD-L1 productions which are necessary for proliferation, cells modulations and T-cells modulations. Only Synthetase enzyme in OPA1 mitochondrial membrane are having the ability of hydrolysis biological molecules, inflammations and phospholipid membranes in vivo ,but
normally followed by the effects of synthase for moderate gamma subunits for producing PLCγ2 which will be moderated by phospholipase effects for PLC alpha production, but in deficiency in the synthase activities or in presence of mutated S6K the osteoblast will be activated, where osteoblast activity is characterized by proper availabilities of S6K, synthase activities, and PLCγ2 synthesis.

Some PLCs isoforms synthesis are involved in multiple stages in TLR4 and interferons regulatory factors (IRFs) synthesis [13]. Where it means the involvement of only PLCγ2 in TLR4 synthesis and in promoting IFN-beta productions for modulate anti-inflammatory effectiveness, but PLCγ1 is promoting IFN gamma activities (PLCγ1 \(\rightarrow\) IFN gamma) which responsible for promoting MHCs class-1 and class two then SIRPα1, TLR4 and PDL1 productions for proliferation, bone growth and T-cells modulations. Also the availability of proper S6K1 for PLCγ1 are so necessary for activating IFN-beta and for TLR4. So, proper PLCγ1 can be considered as important tools produced in vivo for activating IFN gamma and vice versa necessary for regulating PLCγ2 upon BTK activity for anti-inflammatory processes which will be upgraded and moderated by phospholipase activities for PLC alpha, SIRPα1 TLR4 and and for PD-L1 productions.

Therefore, PLCγ1 regulate PLCγ2 production which regulated by tyrosine phosphatase receptors and by phospho-tyrosine receptors “PTyr-R” for activating PLCγ2 productions which then regulate PLC-alpha reproduction for bone growth, for B cells maturation, and for promoting anti-inflammatory steps.

Where, PLCγ2 are basically depend on JAK signaling for SH2B adaptor protein “which are a Tyr kinase receptor family” that necessary for BCR mediate B cells maturations [14] phosphotyrosine “PTyr” are necessary for PLCs synthesis, and for SHP1Src homology region 2 domain containing phosphatase 1 for regulating PLCs productions, for stimulate IFNs productions for anti-inflammatory processes and for proliferations, B-cells maturation, and T-cells modulations.

PLCγ1 is associated with numerous inflammatory diseases due to deficiency in synthase (which depend on availability of Ser and Tyr for PLCγ2 productions, that in some diseases the mutation in S6K can be the main for causing those diseases (due to the deficiency in Ser phosphorylation signaling) and in other cases due to deficiency in proline and in Tyrosine hydroponic amino acids, that healthy immune is depending on the productions of PLCγ1 for activating firstly on infections (by Gamma oxidation) which will promote PLCγ2 productions upon the synthase and BTK activities for modulating anti-inflammatory processes, and then will promote proliferations upon PLC alpha productions due to phospholipase regulations.

PLCγ1 recruit to Colony-stimulating factor-1 “CSF-1” is followed by imp stages for producing PLCγ2 which is necessary for activating anti-inflammatory cycles through activating IFNs which re-activate PLCγ2 via upstream of tyrosine kinase :
The PLCγ1 has the specificity toward colony-stimulating factor receptor synthesis (CSF-1) signaling which expressed on the cell surface that can cause the cells to proliferate and differentiate into specific blood cells, and considered as a class III receptor tyrosine kinase that associated with Neuroinflammation, where PLCγ1 is recruited to the CSF-1 receptor following exposure to the cytokine. [15] PLCγ1 specify for recruit to CSF-1 which necessary for promoting PLCγ2 synthesis for firstly re-activating anti-inflammatory steps then followed promoting proliferation steps through activating PLC alpha, SIRPα1, TLR4 and then PD-L1 productions.

CSF-1 is a members of the IL-1 receptor family regulated by Gamma oxidation by PLCγ1 for promoting PLCγ2 synthesis for re-stimulations IFN beta productions for modulating anti-inflammatory cycles and efficiency.

That, CSF1R-expressing cells may play an anti-inflammatory role or a cancer-suppressive role. [16] As PLCγ1 recruiting to CSF-1 for regulating PLCγ2 synthesis so CSF-1 play necessary role in promoting anti-inflammatory processes which regulated firstly by mitochondrial OPA1 enzymes, by proper S6K production, and by PLCγ1 synthesis.

Also, Tripartite motif (TRIM) 22 plays an important role in interferons (IFNs)-mediated antiviral activity and the Induction of TRIM22 by IFN-γ Involves JAK and PC-PLC/PKC. [17] So PLCs synthesis modulate and regulate Tripartite motif (TRIM) 22 too (which has antimicrobial activities) productions through activating IFNs production.

Also, IFN-γ activates PLC-γ2 via an upstream tyrosine kinase to induce activation of PKC-α. [18] that PLCγ2 regulated by PLCγ1 which can promote IFN-gamma production (through feedback) which has a variety of activities including PLCγ2 re-productions upon the necessity regulations of the upstream of tyrosine kinases for re-activating PKC-α.

PLCγ1 recruited to CSF-1 for two pathways activities 1st / re-activating IFNs productions which regulate MHC class1 and class two for modulating cell-surface protein activities, 2nd / activating PLCγ2 for modulating T-cells , where PLCγ1 involved in the production of TRIM22 for mediating antivirus activities and anti-inflammatory processes through reactivating IFNs productions for PLCγ2 synthesis which modulate T-cells and activate bone growth with activating necessary proliferation. And also PLCγ1 promote IFN gamma which regulate MHC-class-I, MHC class-2 synthesis which promote, SIRPα1, TLR4, and PD-L1 synthesis.

Note that the inhibitions of PLCγ2 productions with PLCγ1 productions will lead to osteoclast, but the proper balance of both PLCγ1 and PLCγ2 productions will lead to osteoblast where PLCγ2 are connected to IFNs productions too.

Also, the Colony-stimulating factor-1 “CSF-1” requires PI3-kinase-mediated metabolism for proliferation [19] PLCγ1 recruited to Colony-stimulating Factor 1 “CSF-1” Depending on mTOR-Ser /Thr phosphorylation signaling for p13k and for proper S6K productions.

And, The inhibitions of of fatty acid synthase “FAS” activity by C75 is resulted in down regulation of phospho-AKT. [20] The inhibition in synthase will reflect down regulations in OPA1 activities and anti-inflammatory processes including PLCγ2 re-productions upon the necessity regulations of the upstream of tyrosine kinases for re-activating PKC-α.

PLCγ1 recruited to CSF-1 for two pathways activities 1st / re-activating IFNs productions which regulate MHC class1 and class two for modulating cell-surface protein activities, 2nd / activating PLCγ2 for modulating T-cells , where PLCγ1 involved in the production of TRIM22 for mediating antivirus activities and anti-inflammatory processes through reactivating IFNs productions for PLCγ2 synthesis which modulate T-cells and activate bone growth with activating necessary proliferation. And also PLCγ1 promote IFN gamma which regulate MHC-class-I, MHC class-2 synthesis which promote, SIRPα1, TLR4, and PD-L1 synthesis.

Note that the inhibitions of PLCγ2 productions with PLCγ1 productions will lead to osteoclast, but the proper balance of both PLCγ1 and PLCγ2 productions will lead to osteoblast where PLCγ2 are connected to IFNs productions too.

Also, the Colony-stimulating factor-1 “CSF-1” requires PI3-kinase-mediated metabolism for proliferation [19] PLCγ1 recruited to Colony-stimulating Factor 1 “CSF-1” Depending on mTOR-Ser /Thr phosphorylation signaling for p13k and for proper S6K productions.

The inhibition in synthase will reflect down regulations in OPA1 membrane and therefore Down regulation in p13k Akt and in S6K productions which necessary for ribosomes repair and for OPA1 repair upon GTPase re-synthesis.

PLCγ2 synthesis activate osteoblast but PLCγ1 production with inhibition in PLCγ2 will activate osteoclast (OC) by inhibiting the inositol 1,4,5-trisphosphate- PLCγ1&2 synthesis are re-modulating variety of cellular pathways including osteoclast (OC) differentiation.

Where, PLCγ2 production is important to be in proper balance with PLCγ1 synthesis for running osteoblast and for inhibiting osteoclast, where the increasing in PLCγ1 productions with inhibition in PLCγ2 will activate osteoclast (OC) by inhibiting re-modulating inositol 1,4,5-trisphosphate “which mediate calcium oscillations and the up-regulation of the nuclear
That, inositol 1,4,5-trisphosphate and diacylglycerol productions require phosphoinositide synthase (PIS) for modulating OC differentiation through regulating transient receptor potential (TRP) channels which requires hydrolysis of phosphatidylinositol 4,5-bisphosphate (PIP) resulting in the generation of inositol 1,4,5-trisphosphate (IP3) and diacylglycerol (DAG).

OPA1 synthase is necessary for creating phosphoinositide synthase (PIS) "regulated by proper S6K production which needed for GTPase synthesis which necessary for OPA1 membrane repairs".

Both PLCγ1 and sphosphoinositide synthase (PIS) are imp for promoting PLCγ2 productions which necessary for upregulate phospholipase activity for PLC alpha for proliferations and bone growth, Where, increasing in PLCγ1 "with reduction or inhibitions in PLCγ2 productions will activate osteoclast but the reactivating proper PLCγ2 synthesis will activate osteoblast. PLCγ2, independent of PLCγ1, was required for receptor activator of NF-xB ligand–induced osteoclastogenesis by differentially regulating nuclear factor of activated T cells c1 (NFATc1). [22] proper PLCγ2 Pathway for modulating osteoclastogenesis Processes mediated by modulating T-cells To complete first the construction of anti-inflammations and the protection followed by the process of building bones growth and cells proliferation in the safety and protection of T-cells and macrophages.

BTK regulate PLCγ2 synthesis which regulate both BCR and Thromboxane-A 2 synthesis, where, CLL disease due to full inhibition in PLCγ2:

Phospholipase Cy2 is Critical for Dectin-1 mediated Ca2+ Flux and Cytokine Production in Dendritic Cells [23].

PLCγ2 has a critical activity in dendritic cells, where is having a Critical function for Development of a Murine Model of Inflammatory Arthritis. [24]

And, as PLCγ2 has a critical activity in dendritic cells for activating NF-xB ligand–induced osteoclastogenesis by differentially regulating nuclear factor-activated T cells c1 "NFATc1".

As PLCγ2 production module first the capacity of T-cells of dendritic cells.

PLCγ2 is critical for B-cell receptor (BCR) for B cells maturation and functions, and PLCγ2 participate in TCR signal transduction and plays a role in T-cell selection [25].

It has been reported that Properdin and factor H production by human dendritic cells modulates their T cell stimulatory. [26] Properdin is plasma glycoprotein that when activated by PLCγ1 (and synthetase) that will be modulated by change unnecessary purines to pyrimidines for rebuilding necessary Tyr, Ser, Pro, then will be directed to x chromosome for translations and purification for being build by identical necessary sequences for being contain identical six thrombospondin that will be ready to be regulated and modulated by PLCγ2 for TXA2 synthesis and for modulating T-cells which mediate cellular and bone growth. The increasing in PLCγ1 productions with deficiency or mutation in S6K and thus in Properdin will inhibit PLCγ2 functions and will reflect decreasing in B cells maturation with decreasing or mutations in the thrombospondin lead to inhibition in TXA2 synthesis and can lead to Autoinflammation and immune dysregulation (APLAIMD) which can cause rare monogenic autoimmune inflammatory disease.

That, the diverse pathologies associated with PLCγ2 are exemplified by distinct genetic variants, where inherited mutations at this locus cause PLCγ2-associated antibody deficiency and immune dysregulation. [27]

Thrombine activation is highly reactivate intermediate the true fibrin monomer and it rapidly, and irreversibly. [28]

That Thrombine is activated by PLCγ2 which intermediate fibrin monomer.

Where, PLCγ2 involved with fibrin formation, where Bruton tyrosine kinase (Btk) activates PLCγ2 ,11,12 leading to thromboxane A2 (TXA2) synthesis. [29]

So, proper PLCγ2 synthesis depend on PLCγ1 and on BTK activities that are necessary for regulating thromboxane-A 2 and fibrin and for re-modulating immune and T cells activities.

Also, the antiplatelet and anti-thrombotic effects of Fe are carried out through oppression of PLCγ2 and subsequent DAG-PKC-TXA2 and IP3-[Ca2+]. [30]

The activation of PLCβ through Gq, which results in the formation of IP3 and diacyl glycerol, plays an important role in mediating αIbβ3 activation. [31]

So in brief the proper S6K, PLCγ1, and BTK necessary for PLCγ2 productions which is necessary for B-cell maturation and T-cells modulations, and necessary for regulating thromboxane-A synthesis. Chronic lymphocytic leukemia [CLL] reflect Inhibition in BTK and in PLCγ2 synthesis which reflect Inhibition or impair in Thromboxane-A:

Proline amino acids are required for Collagen synthesis [32] where, Collagen binds to its receptors and activate both the PLCγ2-DAG-PKC and PI3 kinase/Akt-p38 MAPK cascades, where p38 MAPK can activate cPLA2, which catalyzes arachidonic acid (AA) release to produce thromboxane A2 (TXA 2 ) formation [33].

Bruton’s tyrosine kinase “BTK” activates PLCγ 2 variants mediating ibrutinib resistance in human CLL . [34]

BTK inhibitors [ibrutinib , CNX-774 ] significantly attenuated TPA-induced cell invasion and migration in MCF-7 cells and inhibit the activation of the phospholipase Cy2/PKCb signaling pathways [35].

BTK was initially shown to be defective in the primary immuno-deficiency X-linked a gamma-globulinemia (XLA) and is essential both for B cell development and function of mature. [36]

So, both of Collagen synthesis and BTK are the main functions for re-activating PLCγ2 which catalyzes arachidonic acid (AA) release to produce thromboxane-A2 (TXA 2 ) formation (note the inhibition or mutation in BTK and PLCγ2 will inhibit TXA2 synthesis and will cause Chronic lymphocytic leukemia ), where both BTK and PLCγ2 are so necessary for B cells maturation and are critical for B-cell receptor (BCR), where, inhibition or reduction in BTK and in PLCγ2 will reflect Inhibition in B-cells maturation, inhibition in T-cells modulations, and inhibitions in TXA2 synthesis and will be the result of Chronic lymphocytic leukemia “CLL” disease.

Vascular endothelial growth factor receptor (VEGFR) but not KIT, platelet-derived growth factor receptor (PDGFR) and FMS-like tyrosine kinase 3 (FLT3) are critical for CLL cell viability. [37]

MTOR Ser Thr phosphorylation pathway regulate S6K production and promote VEGF activities for reproducing TXA2 (but through PLCγ2 regulations) in one pathway, and the other pathway is stimulating the PLCγ1 productions and promoting BTK activities for activating PLCγ2 productions which will reactivate the proper TXA2 synthesis and mediate the activities of VEGF for producing TXA2, for reactivating tropomyosin,
and reactivating G-actin filaments activities.

My note, is the synthesis of proper TXA2 in vivo are fully depending on PLCγ2 and consequently on S6K and BTK activities and functions, but only VEGF are not enough and not satisfied for TXA2 synthesis. The proper S6K synthesis which will reactivation the PLCγ1 and DTK which will promote the PLCγ2 synthesis which I can consider it as the main necessary proper tools for TXA2 synthesis for blood synthesis, for bones maturation and for cells growth and then CLL cell viability.

So, PLCγ2 (which basically regulated by ribosomes, by S6K, and by PLCγ1) promote TXA2 synthesis which can stimulate and reactivating VEGF synthesis upon feedback for tropomyosine and for G-actin filaments reactivations for running full cellular Biosynthesis, for blood filtering in veins, and for cellular metabolism.

Chronic lymphocytic leukaemia (CLL) is a malignancy of CD5+ B cells that is characterized by the accumulation of small, mature-appearing lymphocytes in the blood, in bone marrow and in lymphoid tissues due to PLCγ2 inhibition may due to full mutated S6K production.

PLCγ2 synthesis occurred mainly in bone marrow where normal blood synthesis is regulated by skeletal tissue that is having orders from basic ribosomes ,but mature CLL blood are activated and formed only by the activities of mTOR Ser/ Thr signalling which promote the VEGF, tropomyosine synthesis (where both cannot promote TXA2 synthesis without PLCγ2 availability ) that both VEGF and tropomyosine are necessary for reactivate G-actin filaments and re-purify blood in veins.

So why VEGF +tropomyosine is producing white mature cells? VEGF can not regulate directly the PLCγ2 synthesis and consequently can’t regulate TXA2 synthesis but TXA2 synthesis can not be done without PLCγ2 regulations .

Where VEGF responsible for increasing the plasma long lived-plasma cells (LLPC), then the generation of antigen-specific antibody for Durable humoral immunity (which produced by non-proliferating bone marrow. [38] Old blood cells when passes through spleen will be broken to save iron which bind to PLCγ2 for regenerate new blood cells by PLCγ2 which extracted in spleen which are responsible for metals transportations and proliferation for new cells, but inhibition in PLCγ2 with increasing in the mutated S6K will inhibit TXA2 synthesis and will increase long lived plasma which increased by increasing in nutrients-mTOR signalling. The B cell receptor (BCR) (signaling pathway (which regulated by PLCγ2 synthesis and activities) has critical cell survival implications in B-cells malignancies, such as chronic lymphocytic leukaemia (CLL). small molecule tyrosine kinase inhibitors of members of the BCR signaling pathway have proven to be transformational in treatment of CLL. [39] The B-cell receptor (BCR) is a key survival molecule for normal B cells and for most B-cell malignancies.

In CLL, engagement of the BCR (which regulated by PLCγ2) by antigen occurs in vivo, leading to down-regulated expression and to an unanticipated modulation of glycosylation of surface IgM. [40] So inhibition in PLCγ2 synthesis will inhibit BCR signalling function that will lead to inhibition in modulation in IgM which normally done by BCR function for activating B-cells maturation.

IgM autoantibodies, and the evidence that these anti-apoptotic cell IgM natural antibodies can regulate inflammatory responses through ancient pathways of the innate immune system that first arose long before the initial emergence of the adaptive immune system.

My note, PLCγ2 first regulate BCR activities which regulate both IgM and IgD synthesis through synthase enzyme regulation, where IgM is more active and less stable than IgD that IgM necessary for modulating and regulating inflammatory immune response and antiinflammatory processes through modulating T-cells reactivity.

**Results and Conclusion :**

Chronic lymphocytic leukemia [CLL] reflect Inhibition in PLCγ2 synthesis “may due to inhibition in OPA1 synthase” lead to inhibition in CXCR12 where CXCR12 is the main activator and regulator for CXCR4 synthesis Upton phospholipase effects on CXCR12.

Also inhibition in PLCγ2 Bio-Synthesis will reflect reduction or inhibition in thromboxane-A production.

Osteoarthritis “OA” is characterized by a sharp expression in Gamma-Phospholipase C-1 “PLCγ1” (which catabolize inflammations) , with decreasing “or inhibition” in PLCγ2 “PLC beta” productions (which necessary for immune modulation, for B-cell maturation and for T-cells modulation and regulate TXA2 synthesis).

The increasing in PLCγ1 with Deficiency in Ser amino acids , and deficiency in proper S6K, with decreasing or inhibition in OPA1-synthase activity will lead to inhibition in PLCγ2 which lead to diabetes and early Osteoarthritis”OA” prognosis .

PLCγ2 are so necessary for re-modulating T-cells and immune efficiencies, and necessary for regulating antigen and thromboxane-A synthesis.

The inhibitions or reduction or mutations in BTK and in its main proper PLCγ2 productions will cause an inherent inhibition or reduction in CXCL12 then will be followed by inhibition or reduction in CXCR4 then will lead to inhibition in the regulation of B-cell maturation , migration, adhesion, and also lead to severe decreasing in anti-inflammatory processes of immune productive efficiency.

Also inhibition in BTK and PLCγ2 mainly will reflect Inhibition in the two antigens IgM in and IgD synthesis.

Chronic lymphocytic leukemia “CLL” reflect decreasing or inhibition on growth-promoting signaling via the B-cell receptor. The Bruton tyrosine kinase (BTK) is the important for PLCγ2 systems which is necessary for B-cell activities and T-cells modulation.

Bruton tyrosine kinase (Btk) necessary to activates PLCγ2,11,12 which necessary to activate thromboxane A2 and necessary for modulating immune activities and T-cells too.

Both Collagen and BTK pathways are necessary tools for re-activating PLCγ2 which catalyzes arachidonic acid (AA) release to produce thromboxane-A2 (TXA 2 ) synthesis , and necessary for B cells maturation and critical for B-cell receptor (BCR), where, inhibition in BTK and in PLCγ2 will reflect diabetes, Osteoarthritis, and the Chronic lymphocytic leukemia “CLL” disease depending on the percentage of Ser & hydroponic amino acids shortage and depending on the percentage of inhibition of necessary pathways needed for PLCγ2 synthesis and reactivities.

Also, inhibition in the availability of Ser, Tyr, Leu , Pro with inhibition in necessary hydrophobic amino acids synthesis and in BTK and then in PLCγ2 can lead to Osteosarcoma which is a cancer cases that produces immature bone (due to mutations in PLCγ2 and in TLR4 productions) found at the end of long bones, often around the knee.

Deficiency in proline with inhibition in Ser, Tyr, leu (or mutations
in synthase) and in specific beta-subunits-calcium carrier can reflect mutations in the PLCγ2 (beta subunits) productions due to deficiency in proper beta-oxidation that can lead to deficiency or inhibition in the PLCγ2 and PLC alpha, and in MHC class two, that will lead to deficiency or inhibition “or mutations” in “SIRPα1 and in TLR4, PD-L1 then in PD-L1” lead to isolations to that area (due to precipitation of the un functioned calcium by PLCs) that can lead to mutated immature bone and tissue synthesis.

Conflict Of Interest Statement:
The Author declare that the research work has been conducted in the absence of any commercial or financial relationships, that could be construed as a potential conflict of interest.

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Figure 1 - Osteoarthritis linked with diabetes BTK and PLCg2 regulate thromboxane-A Synthesis where their inhibition or mutation reflect CLL diseases. _Discrimination of PLCg2 pathway for modulating T-cells, B-cells maturation and bones growth._

Figure 2 - PLC-gamma-2 regulate both CXCL 12 and CXCR4 upondruton’s tyrosine Kinases “DTK” phosphorylation.
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