Muscle Recovery Is Highlighted by IR Laser Therapy

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Abstract
Introduction: In sports medicine, laser application has been well-established for the recovery of muscles. The mechanisms by which benefits of this kind of therapy can be studied is molecular research approach. Protein-protein interaction network analysis as one of the important complementary studies of proteomics can accelerate this goal by the identification of novel contributing markers.

Methods: By the use of Cytoscape V3.7.1 and its applications, a network of differential expressed proteins (DEPs) from IR laser treatment samples were constructed and analyzed. Six hub-bottlenecks were determined, 4 of which were from differentially expressed proteins.

Results: ClueGO discovered 4 biological processes related to these hub-bottlenecks that their function could alter due to IR laser therapy.

Conclusion: In fact, by the expression changes of hub-bottlenecks including the up-regulation of HSP90s, one of the prominent biological processes in muscle recovery could be activated. This process is called nitric oxide synthase (NOS) activation that could be proposed as one of the underlying mechanisms of IR laser treatments in muscle recovery.

Keywords: Infrared (IR) laser; Myoblast; Proteomics; Protein-protein interaction network analysis

Introduction
Laser therapy has been vastly employed in clinical fields for many years and the subjects including skincare,¹ surgical methods,² oncology,³ and sports medicine.⁴ There are different lines of laser procedures, one of which is infrared laser treatment. The IR laser method has been used for pain-relieving,⁵ dental health,⁶ and sports medicine.⁴

Heavy physical activities could cause some abnormalities in muscles.⁴ As already mentioned, the laser could be beneficial in muscle improvement in a way that the radiation stimulates the cure of wounds and tissues of the traumatized muscles⁴ by the reduction of inflammation.⁷ Molecular investigations including genomics, proteomics, and metabolomics could be helpful to better understand the mechanisms by which the laser could be beneficial in different medical areas.⁸-¹⁰ The effect of laser therapy and non-ionized radiation on gene expression has also been reported for Interleukin 6, vascular endothelial growth factor, MyoD and myogenin expressions in a tissue repairing process.³,¹¹ Other molecular levels such as proteome evaluations could provide key information regarding the molecular mechanisms related to the bio-stimulating effect of laser exposure on protein molecules.⁷ Additional information in this relationship could be achieved via the bioinformatic study of molecular data such as proteomics.¹² Moreover, bioinformatics is a useful method to study mechanism of laser therapy via analyzing the most central biomarkers in an interaction network.⁸ Here, the proteomics data of IR laser treatment on myoblast (C2C12 cell line) of Murine⁴ were employed for our bioinformatic assessments. The protein-protein interaction network analysis of differential expressed proteins (DEPs) obtained from the proteomics investigation was screened for introducing the most essentially contributing biomarkers of the IR Laser mechanism of effects in muscle repair.

Methods

In this study, a list of DEPs including 40 spots from "Effect of IR laser on myoblasts: a proteomic study" was selected. In the original study, a proteomic evaluation of dual laser InfraRed treatment of the C2C12 cell line and the skeletal muscle cell, from Murine was handled. Murine satellite cells were the source of the C2C12 cell line. A multiwave locked system laser (ASA Srl, Vicenza, Italy) was applied; the applied IR laser wavelength was 808 and 905 nm. The main radiation source with average power up to 1.1 W including two synchronized sources (laser diodes) was applied. The first source is a pulsed laser diode that emits 905 nm wavelength and a peak optical power of 25 W. The second one was a laser diode (808 nm) which operated in a continuous mode (power 1.1 W) or a pulsed mode, with the mean optical power output of 550 mW.

The plate containing the C2C12 skeletal muscle cell line was exposed to laser beams with the following scan parameters: 5.6 cms⁻¹, each scan time of 20 seconds, and a total scan time of 8 minutes. The treatment was done under sterile conditions once a day for 3 consecutive days.

The regeneration and repair of muscle cells are assigned with these types of cells. The micrographs of the cell line before and after treatment are presented in the original research. As mentioned earlier, a list of 40 DEPs obtained from IR laser treatments was applied to our bioinformatic approach, the protein-protein interaction network analysis. For this aim, Cytoscape 3.7.1 constructed and analyzed the interaction network via the integrated plugins. Proteins and their interaction queries were dealt with through the use of the STRING db application with the confidence cut-off score of 0.4 as the default selection. Four sources are provided through a STRING platform, including STRING PubMed, STRING Protein, STRING Diseases, and STITCH. The protein queried option was used for the protein-protein interaction network construction. The Uniprot action numbers of 40 proteins were searched against String db. The network was then analyzed for centrality properties including degree (K) and betweenness centrality (BC) as the most important parameters via the Network Analyzer. Proteins with the highest values of degree were called hubs; similarly, nodes with the highest amount of BC were named as bottlenecks. Those proteins with the highest scores of K and BC were called hub-bottlenecks. Hub-bottlenecks were then studied in terms of enrichment properties. In general, genes and proteins can be assessed for their ontology characteristics, including the biological process (BP), the cellular component (CC), and the molecular function (MF). ClueGO and CluePedia identified the BP related to the Hub-bottlenecks with the following statistical criteria: the grouping cut-off score = 0.5 and gene ontology (GO) terms; min. number of genes = 1 and min. gene percentage in term = 3. The Bonferroni step down was the correction method for P value, and the two-sided hypergeometric test was the enrichment/depletion.

Results

Two networks of interaction were constructed in this work, the first one without any additives and the second one with additional neighbors of 50 (Figures 1 and 2).

In the first network of 40 DEPs, there were 40 nodes with 108 interactions. Among these nodes, only 30 were connected. The other DEPs remained as individual nodes. The second network consisted of the queried proteins plus the 50 neighbors, which was 90 nodes in total. This network had 1154 connections. Three of the queried proteins remained as individual nodes. These nodes were Clec3b, Nlrp10, and Phldb2 that did not come to the connection with other nodes. Further analysis based on network centrality showed that some of the proteins possessed valuable properties in the strength of the network. In Table 1, the list of the proteins with the highest centrality scores are gathered and ranked based on the degree value.
The centrality analysis of the second network showed that most of the hub-bottleneck proteins were from query DEPs. Only two were from neighbor proteins, which were Gapdh and Itpkb. Actb, the most highlighted hub-bottleneck, was with a degree of 65 and betweenness centrality of 0.06.

Biological processes related to the central nodes were provided by ClueGO. The terms are clustered and the identified groups are represented based on significant parameters in Figure 3.

Nitric oxide synthase (NOS) regulator activity was the leading biological process group with about 48% of the entire BPs in this query in ClueGO. The presence of 2 proteins per term was obtained for the hub-bottleneck DEPs. Likewise, HSP90aa1 and HSP90ab1 belonged to the NOS regulator activity; Actg1 and Actb were the related structural constituent of the postsynaptic actin cytoskeleton. The other 2 nodes which were not from differentially expressed proteins were not common in a term; however, Itpkb and Gapdh were linked to inositol-1,4,5-trisphosphate 3-kinase activity and Peptidyl-cysteine 5-nitrosylase activity individually.

**Discussion**

Bioinformatics including protein-protein interaction network analysis could help in the light of understanding the mechanism in which the effect of applied treatment on living organisms is detected. Laser therapy, which has been used since the 1970s, is one of the common methods in muscle recovery treatments. Here, to better recognize the IR effectiveness in muscle cells, the DEPs from the original proteomics study of IR Laser treatment on myoblast cell were evaluated in silico. Two PPI networks were constructed from the merely queried proteins (DEPs) and the queried individuals plus 50 neighbor proteins. The condense interaction between these proteins and the other nodes indicated their important interaction values. Moreover, the first network query without any additives showed that the DEPs were in a cluster of interactions. The network centrality analysis implied the presence of essential nodes that were vital for network stability. These nodes were recognized as 6 central proteins in the network of IR laser treatment on the C2C12 cell line.

Among them, Actb, Hsp90aa1, Hsp90ab1, and Actg1 were the DE hub-bottlenecks. Gapdh and Itpkb were the other 2 hub-bottlenecks, but they were from the neighbor proteins. There were four biological processes linked to the DE central proteins and accordingly, NOS regulator activity and the structural constituent of the postsynaptic actin cytoskeleton could definitely get affected as a result of dysregulation of DE hub-bottlenecks.

Chaperons are protein molecules that bind to the non-native proteins to obtain biological function. These molecular machines are overexpressed in the stress condition and protect the body. Among them, Actb, Hsp90aa1, Hsp90ab1, and Actg1 were the DE hub-bottlenecks. Gapdh and Itpkb were the other 2 hub-bottlenecks, but they were from the neighbor proteins. There were four biological processes linked to the DE central proteins and accordingly, NOS regulator activity and the structural constituent of the postsynaptic actin cytoskeleton could definitely get affected as a result of dysregulation of DE hub-bottlenecks.

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The well-known molecular mechanism of muscle contraction is tied to the interaction between actin and myosin filaments. Different variants of actin molecules, including alpha-actin, beta-actin, and gamma actin, are...
identified and involved in muscle contraction and also other vital cellular functions such as proliferation and synthetic or secretory activities. Since the overexpression of Actb and actg1 (the other 50% of queried DEPs) is associated with the repair procedure in muscle recovery, highlighting these two variants of actin via network analysis reveals their important role by laser radiation. It can be concluded that the IR laser can be considered as repair promotion and cell protection in different muscle stress conditions. The expression change of actin after low-intensity IRs radiation in skin and muscle is reported by Fonseca et al. In this document, it is reported that actin mRNA expression changes, depending on fluency, frequency and time of its exposure to radiation in the skin and muscle of Wistar rats. Although the expression changes of the other 2 hub-bottlenecks are not reported in the original study, they might have some roles in the underlying laser molecular mechanism. Glyceraldehyde-3-phosphate dehydrogenase, Gapdh, is known as a metabolic enzyme that is involved in sugar metabolism. The role of Gapdh in maintaining adequate glycolytic flux and related pathways are discussed by Seidler. Its character in muscle recovery and its function in stress condition and also after laser therapy have been reported by many studies. Its participation as a central protein in this study refers to a possible critical role of the IR laser in the evocation of cell metabolism. Itpkb, inositol trisphosphate 3-kinase B, as another neighbor hub-bottleneck in the PPI network, has been shown as a pathogenic factor for Alzheimer disease by increasing the level of amyloid beta in patients. Therefore, as a novel central protein in our network, it is vital to assess its feasible alterations as a possible side effect of laser treatments. Altogether, the central nodes and their corresponding biological processes could pinpoint the molecular mechanism of IR laser effectiveness.

Conclusion
IR Laser therapy is an effective method to promote muscle repairing by influencing central proteins of the PPI network of the C2C12 Cell Line. One of the crucial mechanisms could be through the activation of NOS. The Gapdh and Itpkb, regardless of not being DEPS, could also show potential in laser treatments; nevertheless, an intensive investigation is required for this claim.

Ethical Considerations
Not applicable.

Conflict of Interests
The authors declare no conflict of interest.

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References
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