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Original Scientific Paper

SIGNIFICANCE OF FIBROBLAST GROWTH FACTOR 23 IN ACUTE KIDNEY INJURY

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SUMMARY – Acute kidney injury is a clinical syndrome associated with increased patient morbidity and mortality, as well as serious short-term and long-term consequences, especially in the perioperative period. Yet, patients having suffering from temporary acute kidney injury and achieving full recovery of kidney function usually complain of poor quality of life associated with loss of energy and limited physical activity. Therefore, there is a necessity for a novel biomarker of acute kidney injury with better features than currently used serum creatinine and urine output. So far, several investigations have demonstrated that the fibroblast growth factor 23 could be that desperately searched novel biomarker of acute kidney injury. It cannot only detect kidney dysfunction at the time but also before the injury process begins. Moreover, serum levels of the fibroblast growth factor 23 can predict adverse progression of the kidney injury. However, the role of the fibroblast growth factor 23 in the acute but also in chronic kidney dysfunction is still a riddle that requires additional research to clarify it.

Key words: Acute kidney injury; Biological biomarkers; Fibroblast growth factor 23; Perioperative period

Acute Kidney Injury

Acute kidney injury (AKI) is a frequent perioperative complication and when it occurs independently, it increases patient morbidity, mortality and poor outcome¹⁻⁴. AKI is a clinical syndrome characterized by a sudden or persistent decrease in kidney function⁵⁻⁹. Due to reduction in the glomerular filtration rate (GFR), it is usually manifested as an increase in serum creatinine or decrease in urine output. Although GFR is the accepted indicator of renal function, due to its difficult measurement it is usually assessed from

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serum creatinine levels and urine output¹⁰. However, elevation of serum creatinine and decrease in urine output do not allow clinicians to make a timely diagnosis of AKI, or to detect the site or severity of injury¹¹. Serum creatinine is a renal biomarker the values of which vary widely with the patient clinical condition and therapy; thus, elevation in serum creatinine is usually delayed by several hours and occasionally by several days until reaching a steady value¹². Also, about 50% of renal function must be lost prior to serum creatinine starting to rise¹². On the other hand, diuresis is not a sensitive renal biomarker because many hospitalized patients, especially critically ill, receive diuretics in their daily therapy, so oliguria cannot be detected¹². In addition, great attention should be paid to surgical patients that usually develop temporary oliguria postoperatively due to perioperative multifactorial and complex hemodynamic and volume

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disturbances, which lead to hypovolemia and subsequent renal hypoperfusion¹³⁻¹⁶. It is considered as a normal physiological renal response, so a clinician has to distinguish it from AKI.

Kidney is a robust organ and can put up with exposure to many noxious impacts without any significant structural and functional renal impairment¹⁰. However, when this renal impairment does occur, it usually implies that severe multiorgan derangements have happened and that the patient's condition is rather serious. Every clinician should be aware of a timely narrow therapeutic window of 24-48 hours between kidney submission to the high risk and noxious factors, and AKI genesis. In this important period, preventive and supportive measures must be undertaken and a novel biomarker of AKI should demonstrate its sensitivity and specificity. Therefore, small changes in serum creatinine values and urine output are independently associated with mortality and should be early detected and alert every clinician to start appropriate patient monitoring and therapy¹⁷⁻²⁰. Many studies have shown that even acute and mild kidney dysfunction can have severe sequels⁵⁻⁹. Moreover, 30% of patients who have experienced temporary uncomplicated AKI with full recovery of kidney function are at an increased risk of chronic kidney disease (CKD) development19,20,21.

The etiology and pathophysiology of AKI are complex and multifactorial²²⁻²⁴. Nevertheless, we can conclude that the major kidney risk factors are hypotension, ischemia and reperfusion, inflammation and nephrotoxins^{23,24}. Their detrimental impact on the kidney function and structure can be reversible, especially if therapeutic measures are applied in the narrow therapeutic renal window. Thereby, the main goal is to detect every insult to the kidney as early as possible. Therefore, a novel AKI biomarker is required. Till then, AKI should be always if possible predicted by the clinician and a supportive and nephroprotective strategy should be initiated¹³. These measures consist of optimization of hemodynamic parameters like maintaining systolic pressure >80 mm Hg, mean arterial pressure >65 mm Hg and renal perfusion pressure¹³. However, we consider that optimization of hemodynamic parameters is best managed by timely and early goal directed therapy using less invasive hemodynamic monitoring if possible¹³. Another puzzle in daily management of AKI is when to initiate the supportive renal replacement therapy. Nowadays, there is still a tendency to postpone renal replacement therapy as long as possible¹⁰. Renal replacement therapy deferring is usually justified by the patient hemodynamic instability, dialysis catheter associated complications, and activation of the patient immune and inflammation system due to bioincompatible dialysis membrane^{25,26}. However, studies have demonstrated that late renal replacement therapy initiation is connected with increased patient mortality, longer duration of renal replacement therapy and hospital stay, compromised recovery of renal function, and in the end with a greater risk of progression to CKD and dialysis dependence²⁷⁻³¹. On the other hand, early initiation of renal replacement therapy as soon as the patient develops stage 3 of AKI according to the RIFLE criteria is warranted by the benefit of avoiding exposure to the deleterious effects of metabolic waste products and abnormalities, or volume overload¹⁰. The major goal of the novel kidney biomarker would be to predict renal recovery and to give clear answer to the relevant question when to initiate and discontinue renal replacement therapy¹⁰. By then, renal replacement therapy initiation will remain a clinical risk-benefit ratio decision based on the already developed complications manifested in the fluid, metabolic and electrolyte system of the individual patient¹⁰.

Thus, the desperately needed novel kidney biomarker should have the ability to early diagnose and predict the AKI course, identify the etiology, location and severity of renal injury, provide individual patient risk stratification, define timing of renal replacement therapy initiation and discontinuation, and to be cost-effective, reliable and easy to use in clinical practice^{11,32,33}.

Fibroblast Growth Factor 23

Fibroblast growth factor 23 (FGF23) is a bone produced novel hormone that for now has excellent features to be considered a new and early AKI biomarker³⁴. It is generated and secreted primarily in bones by osteoblasts and osteocytes³⁵. It pertains to the FGF19 subfamily and acts as an endocrine phosphaturic hormone, i.e. phosphatonine³⁶. So, the major physiological role of FGF23 is to maintain phosphate homeostasis³⁷. Preserving the systemic phosphate ho-

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meostasis is essential for the body. The reason is that various cellular functions ranging from cellular signaling to enzymatic activities depend on the phosphate balance³⁸. When this balance is disrupted, severe vital organ system malfunctions arise³⁸. Kidney is the organ responsible for preserving phosphate homeostasis by a closed negative feedback loop between the kidneys, bones and parathyroid glands. When this loop becomes open, serum phosphate levels start to increase. So, hyperphosphatemia increases parathyroid hormone (PTH) and FGF23 secretion, which in turn enhance phosphaturia in a similar manner. As a result of kidney dysfunction, this negative feedback system is disrupted and phosphate serum levels augmentation continues. The only difference between PTH and FGF23 is that PTH increases phosphaturia in a few hours, whereas FGF23 reacts after several days^{39,40}. The main phosphaturic mechanism of FGF23 is performed by inhibiting renal tubular reabsorption of phosphate by reducing the number of sodium-dependent phosphate IIa and IIc cotransporters in the apical brush border of the renal proximal tubule⁴¹⁻⁴³. Everything we currently know about the FGF23 intracellular mechanism of action on renal phosphate excretion is that it has to connect with its main renal specific receptor FGFR1 together with the help of the Klotho transmembrane protein as a coreceptor to accomplish the effect^{41,42}. Hence, the Klotho protein is the relevant factor that determines the FGF23 effects on target organs such as the kidney, parathyroid glands, pituitary gland and choroid plexus in the brain⁴⁴. Moreover, other avenues of maintaining phosphate homeostasis are decreasing intestinal phosphate absorption through direct action of reducing the number of sodium-dependent phosphate IIb cotransporters in the gastrointestinal mucosal apical border, and indirectly through blocking calcitriol synthesis by reducing renal 1-alfa-hydroxylase activity and enhancing calcitriol catabolism by stimulating 24-alfa hydroxylase activity^{34,36,45}. The sequel of this avenue is diminished vitamin D-dependent intestinal phosphate absorption⁴⁵. However, the FGF23 fashion of degradation is not well understood⁴⁶. According to the literature, there are two forms of circulating FGF23 in the serum. One is intact and full-length form of the FGF23 which has biological activity and the other is inactive C-terminal fragment of the

FGF23 without biological activity in the serum³⁶. The assessed half-life of the intact FGF23 in the circulation is about 58 minutes⁴⁷.

Fibroblast Growth Factor 23 and Chronic Kidney Disease

Studies of the FGF23 in patients suffering from CKD have revealed that the capital drivers for the increased FGF23 production are intestinal phosphate absorption after high oral phosphate entry, calcitriol, and PTH directly and indirectly by increased 1,25dihydroxy vitamin D3-dependent production^{36,48-51}. Thus, FGF23 serum levels already significantly start to rise during early stages of CKD before serum phosphate derangements can be detected^{46,52,53}. Thereby, in patients with end-stage CKD, they are hundredsto thousands-fold higher than normal values^{46,52,53}. Hence, we conclude that FGF23 could be the early biomarker of renal deterioration in the progress of CKD, and its serum values can predict the CKD outcome³⁶. Also, FGF23 increased serum values are independently associated with higher mortality, therapy resistant secondary hyperparathyroidism, left ventricular hypertrophy and cardiovascular mortality in CKD patients^{38,52}. This adverse effect of elevated FGF23 serum concentrations on cardiovascular system is not well understood. Some evidence supports direct FGF23 toxicity on the cardiac and vascular endothelial function through low-affinity Klotho independent FGF receptors^{54,55}. Other evidence demonstrates the FGF23 cardiovascular toxicity due to disruption in the calcium-phosphate metabolism homeostasis³⁶. However, regardless of the mechanism of action, the result is the same.

Fibroblast Growth Factor 23 and Acute Kidney Injury

There are not so many studies on the role of FGF23 in AKI, but recent studies have revealed that AKI is associated with raised serum levels of FGF23⁵⁶⁻⁶⁰. In addition, studies showed that elevated serum levels of FGF23 in AKI patients were connected with unfavorable outcome in terms of a significantly increased risk of death or need of renal replacement therapy^{57,58}. Serum levels of FGF23 in AKI rise independently of the already mentioned drivers in CKD patients such as high

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oral phosphate entry, calcitriol and PTH⁵⁷. However, the precise mechanism of the increased FGF23 levels has not yet been elucidated. Increased production rather than decreased elimination of FGF23 contributes to elevated levels of FGF23 in AKI, so impaired renal FGF23 scavenging due to reduction of the glomerular function units is not the major pathway for raising the FGF23 levels⁶¹. Also, another pathway that can contribute to FGF23 serum elevation may be deficiency of the important FGF23 coreceptor Klotho in the injured kidney⁶². The Klotho deficiency leads to the FGF23 resistance resulting in raised FGF23 serum levels. Besides, enhanced bone production of FGF23 in AKI patients was demonstrated by immunohistochemical and Western blot analyses of the bones also as a mechanism contributing to elevated FGF23 serum levels^{63,64}. Yet, the responsible kidney driven factor for increased FGF23 bone production and the actual pathway of interaction in this novel bone-kidney axis still remains obscure³⁶. In addition, due to many difficulties in detecting AKI in perioperative settings, FGF23 manifested as a reliable biomarker of the acute kidney dysfunction since it begins to rise significantly within 24 hours postoperatively⁵⁷. Furthermore, one pilot study in pediatric population submitted to cardiac surgery indicated that elevated preoperative FGF23 serum levels were also associated with a two-fold increased risk of AKI development, yet correlating with positive fluid balance, length of mechanical ventilation, and length of stay at the intensive care unit⁵⁷.

Conclusion

In conclusion, the currently used kidney injury biomarkers cannot detect AKI timely and usually their concentrations start to rise when the injury process is in the late phase. The goal of the new and early kidney injury biomarker is to predict and diagnose AKI with the aim to start the nephroprotective treatment before the damage has happened. FGF23 could be a novel promising biomarker of AKI, especially in the complex perioperative period. However, additional research in the field is needed.

References

 Doi K, Negishi K, Ishizu T, Katagiri D, Fujita T, Matsubara T, et al. Evaluation of new acute kidney injury biomarkers in a mixed intensive care unit. Crit Care Med. 2011;39:1-6.

- Abelha FJ, Botelho M, Fernandes V, Barros H. Determinants of postoperative acute kidney injury. Crit Care. 2009;13:R79.
- Lassnigg A, Schmid ER, Hiesmayr M, Falk C, Druml W, Bauer P. Impact of minimal increases in serum creatinine on outcome in patients after cardiothoracic surgery: do we have to revise current definitions of acute renal failure? Crit Care Med. 2008;36:1129-37.
- Bihorac A, Yavas S, Subbiah S, Hobson CE, Schold JD, Gabrielli A, et al. Long-term risk of mortality and acute kidney injury during hospitalization after major surgery. Ann Surg. 2009;249:851-8.
- Chertow GM, Burdick E, Honour M, Bonventre JV, Bates DW, et al. Acute kidney injury, mortality, length of stay, and costs in hospitalized patients. J Am Soc Nephrol. 2005;16:3365-70.
- Hoste EA, Clermont G, Kersten A, Venkataraman R, Angus DC, De Bacquer D, et al. RIFLE criteria for acute kidney injury are associated with hospital mortality in critically ill patients: a cohort analysis. Crit Care. 2006;10:R73.
- Lassnigg A, Schmidlin D, Mouhieddine M, Bachmann LM, Druml W, Bauer P, et al. Minimal changes of serum creatinine predict prognosis in patients after cardiothoracic surgery: a prospective cohort study. J Am Soc Nephrol. 2004;15:1597-605
- Levy EM, Viscoli CM, Horwitz RI. The effect of acute renal failure on mortality. A cohort analysis. JAMA. 1996;275:1489-94.
- 9. Uchino S, Bellomo R, Goldsmith D, Bates S, Ronco C, *et al.* An assessment of the RIFLE criteria for acute renal failure in hospitalized patients. Crit Care Med. 2006;34:1913-7.
- Kidney Disease: Improving Global Outcomes (KDIGO) Acute Kidney Injury Work Group. KDIGO Clinical Practice Guideline for Acute Kidney Injury. Kidney Int Suppl. 2012;2:1-138.
- Siew ED, Ware LB, Gebretsadik T, Shintani A, Moon KG, Wickersham N, et al. Urine neutrophil gelatinase-associated lipocalin moderately predicts acute kidney injury in critically ill adults. J Am Soc Nephrol. 2009;20:1823-32. Epub 2009 Jul 23.
- Lamiere N. The diagnostic role of novel biomarkers in acute kidney injury in critically ill patients. Int J Intensive Care. 2011;4:118-21.
- 13. Brienza N, Giglio MT, Dalfino L. Surgical and intensive care strategies to prevent renal failure. Int J Intensive Care. 2011;4:100-8.
- Lobo DN, Macafee DA, Allison SP. How perioperative balance influences postoperative outcomes. Best Pract Res Clin Anaesthesiol. 2006;20:439-55.
- Gann DS, Kenney PR. Special problems of fluid and electrolyte management in surgery. Chan JCM, Gill JR, eds. Kidney Electrolyte Disorders. Churchill Livingstone Inc., USA 1990;343-62.

- Alpert RA, Roizen MF, Hamilton WK, Stoney RJ, Ehrenfeld WK, Poler SM, et al. Intraoperative urinary output does not predict postoperative renal function in patients undergoing abdominal aortic revascularisation. Surgery. 1984;95:707-11
- Metnitz PG, Krenn CG, Steltzer H, Lang T, Ploder J, Lenz K, et al. Effect of acute renal failure requiring renal replacement therapy on outcome in critically ill patients. Crit Care Med. 2002;30:2051-8.
- Levy EM, Viscoli CM, Horwitz RI. The effect of acute renal failure on mortality: a cohort analysis. JAMA. 1996;275:1489-94
- Harel Z, Chan CT. Predicting and preventing acute kidney injury after cardiac surgery. Curr Opin Nephrol Hypertens. 2008;17:624-8.
- Reddy VG. Prevention of postoperative acute renal failure. J Postgrad Med. 2002;48:64-70.
- 21. Venkataraman R. Can we prevent acute kidney injury? Crit Care Med. 2008;36:S166-S171.
- Kellum JA. Acute renal failure, interdisciplinary knowledge and the need for standardization. Curr Opin Crit Care. 2005;11:525-6.
- Kellum JA. Acute kidney injury. Crit Care Med. 2008;36:S141-S145
- Lameire N, Van BW, Vanholder R. Acute kidney injury. Lancet. 2008;372:1863-5.
- 25. Conger JD. Does hemodialysis delay recovery from acute renal failure? Semin Dial. 1990;3:146-8.
- Schortgen F, Soubrier N, Delclaux C, Thuong M, Girov E, Brun-Buisson C, et al. Hemodynamic tolerance of intermittent hemodialysis in critically ill patients: usefulness of practice guidelines. Am J Respir Crit Care Med. 2000;162:197-202.
- 27. Palevsky PM, Baldwin I, Davenport A, Goldstein S, Paganini E, *et al.* Renal replacement therapy and the kidney: minimizing the impact of renal replacement therapy on recovery of acute renal failure. Curr Opin Crit Care. 2005;11:548-54.
- Conger JD. A controlled evaluation of prophylactic dialysis in post-traumatic acute renal failure. J Trauma. 1975;15:1056-63.
- 29. Fischer RP, Griffen WO Jr, Reiser M, Clark DS. Early dialysis in the treatment of acute renal failure. Surg Gynecol Obstet. 1966;123:1019-23.
- Kleinknecht D, Jungers P, Chanard J, Babanel C, Ganeval D. Uremic and non-uremic complications in acute renal failure: evaluation of early and frequent dialysis on prognosis. Kidney Int. 1972;1:190-6.
- Parsons FM, Hobson SM, Blagg CR, McCracken BH. Optimum time for dialysis in acute reversible renal failure. Description and value of an improved dialyser with large surface area. Lancet. 1961;1:129-34.
- Devarajan P. Proteomics for biomarker discovery in acute kidney injury. Semin Nephrol. 2007;27:637-51.

- Devarajan P. Proteomics for the investigation of acute kidney injury. Contrib Nephrol. 2008;160:1-16.
- Quarles LD. Role of FGF23 in vitamin D and phosphate metabolism: implications in chronic kidney disease. Exp Cell Res. 2012;318:1040-8. Epub 2012 Mar 7.
- 35. Riminucci M, Collins MT, Fedarko NS, Cherman N, Corsi A, White KE, *et al.* FGF-23 in fibrous dysplasia of bone and its relationship to renal phosphate wasting. J Clin Invest. 2003;112:683-92.
- Kocełak P, Olszanecka-Glinianowicz M, Chudek J. Fibroblast growth factor 23 – structure, function and role in kidney diseases. Adv Clin Exp Med. 2012;21:391-401.
- 37. Quarles LD. Endocrine functions of bone in mineral metabolism regulation. J Clin Invest. 2008;118:3820-8.
- 38. Russo D, Battaglia Y. Clinical significance of FGF-23 in patients with CKD. Int J Nephrol. 2011;364890:5 pages. Epub 2011 Apr 26.
- Nishida Y, Taketani Y, Yamanaka-Okumura H, Imamura F, Taniquchi A, Sato T, et al. Acute effect of oral phosphate loading on serum fibroblast growth factor 23 levels in healthy men. Kidney Int. 2006;70:2141-7.
- Reiss E, Canterbury J, Bercovitz M, Kaplan EL. The role of phosphate in the secretion of parathyroid hormone in men. J Clin Invest. 1970;49:2146-9.
- 41. Gattineni J, Bates C, Twombley K, Dwarakanath V, Robinson ML, Goetz R, et al. FGF23 decreases renal NaPi-2a and NaPi-2c expression and induces hypophosphatemia in vivo predominantly via FGF receptor 1. Am J Physiol Renal Physiol. 2009;297:F282-F291.
- 42. Liu S, Vierthaler L, Tang W, Zhou J, Quarles LD. FGFR3 and FGFR4 do not mediate renal effects of FGF23. J Am Soc Nephrol. 2008;19:2324-50.
- Baum M, Schiavi S, Dwarakanath V, Quigley R. Effect of fibroblast growth factor-23 on phosphate transport in proximal tubules. Kidney Int. 2005;68:1148-53.
- Urakawa I, Yamazaki Y, Shimada T, Iijima K, Hasegawa H, Okawa K, et al. Klotho converts canonical FGF receptor into a specific receptor for FDF23. Nature. 2006;444:770-4.
- Miyamoto K, Ito M, Kuwahata M, Kato S, Segawa H. Inhibition of intestinal sodium-dependent inorganic phosphate transport by fibroblast growth factor 23. Ther Apher Dial. 2005;9:331-5.
- 46. Larsson T, Nisbeth U, Ljunggren O, Juppner H, Jonsson Kb. Circulating concentration of FGF-23 increases as renal function declines in patients with chronic kidney disease, but does not change in response to variation in phosphate intake in healthy volunteers. Kidney Int. 2003;64:2271-9.
- 47. Khosravi A, Cutler C, Kelly M, Chang R, Royal RE, Sherry RM, *et al.* Determination of the elimination half-life of fibroblast growth factor-23. J Clin Endocrinol Metab. 2007;92:2374-7.
- 48. Burnett SM, Gunawardene SC, Bringhurst FR, Juppner H, Lee H, Finkelstein JS, et al. Regulation of C-terminal and

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- intact FGF-23 by dietary phosphate in men and women. J Bone Miner Res. 2006;21:1187-96.
- 49. Antoniucci DM, Yamashita T, Portale AA. Dietary phosphorus regulates serum fibroblast growth factor-23 concentrations in healthy men. J Clin Endocrinol Metab. 2006;91:3144-9.
- Lavi-Moshayoff V, Wasserman G, Meir T, Silver J, Naveh-Manyi T. PTH increases FGF23 gene expression and mediates the high-FGF23 levels of experimental kidney failure: a bone parathyroid feedback loop. Am J Physiol Renal Physiol. 2010;299:F882-F889.
- 51. Wesseling-Perry K, Pereira RC, Sahney S, Gales B, Wang HJ, Elashoff R, *et al.* Calcitriol and doxercalciferol are equivalent in controlling bone turnover, suppressing parathyroid hormone, and increasing fibroblast growth factor-23 in secondary hyperparathyroidism. Kidney Int. 2011;79:112-9.
- Gutierrez OM, Mannstadt M, Isakova T, Rauch-Hain JA, Tamez H, Shah A, et al. Fibroblast growth factor 23 and mortality among patients undergoing hemodialysis. N Engl J Med. 2008;359:584-92.
- 53. Isakova T, Wahl P, Vargas GS, Gutierrez OM, Scialla J, Xie H, et al. Fibroblast growth factor 23 is elevated before parathyroid hormone and phosphate in chronic kidney disease. Kidney Int. 2011;79:1370-8.
- Mirza M, Larsson A, Lind L, Larsson TE. Circulating fibroblast growth factor-23 is associated with vascular dysfunction in the community. Atherosclerosis. 2009;205:385-90.
- 55. Razzaque M. Does FGF23 toxicity influence the outcome of chronic kidney disease? Nephrol Dial Transplant. 2009;24:4-7.
- Leaf DE, Wolf M, Waikar SS, Chase H, Christov M, Cremers S, et al. FGF-23 levels in patients with AKI and risk of adverse outcomes. Clin J Am Soc Nephrol. 2012;7:1217-23. Epub 2012 Jun 14.

- 57. Christov M, Waikar SS, Pereira RC, Havasi A, Leaf DE, Goltzman D, et al. Plasma FGF23 levels increase rapidly after acute kidney injury. Kidney Int. 2013;84:776-85. Epub 2013 May 8.
- Charnow JA. High FGF-23 in AKI raises death risk. Renal & Urology News. Haymarket Media. 2011. HighBeam Research. 3 Feb 2014. Available at: http://www.highbeam.com.
- 59. Ali FN, Hassinger A, Price H,Langman CB. Preoperative plasma FGF23 levels predict acute kidney injury in children: results of a pilot study. Pediatr Nephrol. 2013;28:959-62. Epub 2013 Jan 13.
- 60. Zhang M, Hsu R, Hsu CY, Kordesch K, Nicasio E, Cortez A, *et al.* FGF-23 and PTH levels in patients with acute kidney injury: a cross-sectional case series study. Ann Intensive Care. 2011;1:21.doi:10.1186/2110-5280-1-21.
- 61. Isakova T, Xie H, Barchi-Chung A, Vargas G, Snowden N, Houston J, *et al.* Fibroblast growth factor 23 in patients undergoing peritoneal dialysis. Clin J Am Soc Nephrol. 2011;6:2688-95.
- 62. Koh N, Fujimori T, Nishiguchi S, Tamori A, Shiomi S, Nakatani T, *et al.* Severely reduced production of klotho in human chronic renal failure kidney. Biochem Biophys Res Commun. 2001;280:1015-20.
- 63. Stubbs JR, He N, Idiculla A, Gillihan R, Liu S, David V, *et al.* Longitudinal evaluation of FGF23 changes and mineral metabolism abnormalities in a mouse model of chronic kidney disease. J Bone Miner Res. 2012;27:38-46.
- 64. Pereira RC, Juppner H, Azucena-Serrano CE, Yadin O, Salusky IB, Wesseling-Perry K, *et al.* Patterns of FGF-23, DMP1, and MEPE expression in patients with chronic kidney disease. Bone. 2009;45:1161-8.

Sažetak

ULOGA ČIMBENIKA RASTA FIBROBLASTA 23 U AKUTNOJ OZLJEDI BUBREGA

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Akutna ozljeda bubrega je klinički sindrom povezan s povišenim rizikom pobola i smrtnosti te kratkotrajnim i dugotrajnim posljedicama, osobito kod bolesnika u perioperacijskom razdoblju. Zbog toga potreban je novi biomarker akutne bubrežne ozljede, bolje specifičnosti i osjetljivosti od serumske vrijednosti kreatinina i diureze. Nekoliko dosadašnjih radova je pokazalo da čimbenik rasta fibroblasta 23 može pravodobno prepozanti poremećaj bubrežne funkcije tijekom samog procesa oštećenja. Također, povišene serumske vrijednosti čimbenika rasta fibroblasta 23 mogu upozoriti na nepovoljan ishod bubrežne ozljede. Međutim, uloga čimbenika rasta fibroblasta 23 u akutnoj bubrežnoj ozljedi je velika nepoznanica te su potrebna daljnja istraživanja.

Ključne riječi: Akutna ozljeda bubrega; Biološki biljezi; Čimbenik rasta fibroblasta 23; Perioperacijsko razdoblje