

## Öğr.Gör. ASLI İNCİ

### Kişisel Bilgiler

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### Eğitim Bilgileri

Lisans Yandal, Gazi Üniversitesi, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, Türkiye 2014 - 2017

### Akademik Unvanlar / Görevler

Öğretim Görevlisi, Gazi Üniversitesi, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2018 - Devam Ediyor

### SCI, SSCI ve AHCI İndekslerine Giren Dergilerde Yayınlanan Makaleler

- I. **Autism: Screening of inborn errors of metabolism and unexpected results**  
İNCİ A., ÖZASLAN A., OKUR İ., BİBEROĞLU G., GÜNEY E., EZGÜ F. S., TÜMER L., İŞERİ E.  
AUTISM RESEARCH, 2021 (SCI İndekslerine Giren Dergi)
- II. **The chemical chaperone 4-phenylbutyrate enhances alpha-galactosidase activity subsequent to stop-codon read-through therapy with triamterene in Fabry R227X fibroblasts**  
DÜNDAR H., BİBEROĞLU G., İNCİ A., IŞIK GÖNÜL İ., OKUR İ., TÜMER L., EZGÜ F. S.  
MOLECULAR GENETICS AND METABOLISM, cilt.132, sa.2, 2021 (SCI İndekslerine Giren Dergi)
- III. **Beneficial Effects of Modified Atkins Diet in Glycogen Storage Disease Type IIIa**  
Olgac A., İNCİ A., OKUR İ., BİBEROĞLU G., Oguz D., EZGÜ F. S., Kasapkara C. S., Aktas E., TÜMER L.  
ANNALS OF NUTRITION AND METABOLISM, cilt.76, sa.4, ss.233-241, 2020 (SCI İndekslerine Giren Dergi)
- IV. **A new NBIA patient from Turkey with homozygous C19ORF12 mutation**  
Kasapkara C. S., TÜMER L., Gregory A., Ezgu F., İNCİ A., Derinkuyu B. E., Fox R., Rogers C., Hayflick S.  
ACTA NEUROLOGICA BELGICA, cilt.119, sa.4, ss.623-625, 2019 (SCI İndekslerine Giren Dergi)
- V. **Vitamin D Levels and Bone Mineral Density in Inborn Errors of Metabolism Requiring Specialised Diets**  
Olgac A., İNCİ A., OKUR İ., Ezgu F., BİBEROĞLU G., Turner L.  
JCPS-P-JOURNAL OF THE COLLEGE OF PHYSICIANS AND SURGEONS PAKISTAN, cilt.29, sa.12, ss.1207-1211, 2019 (SCI İndekslerine Giren Dergi)
- VI. **Patient With Niemann-Pick Type C Presenting With a Jaw Mass Characterized With Lymph Node Involvement by Niemann-Pick Cells**  
İNCİ A., OKUR İ., ESENDAĞLI G., OKUR A., Olgac A., EZGÜ F. S., TÜMER L.  
JOURNAL OF PEDIATRIC HEMATOLOGY ONCOLOGY, cilt.40, sa.3, ss.243-245, 2018 (SCI İndekslerine Giren Dergi)

### Diğer Dergilerde Yayınlanan Makaleler

- I. **Citrullinemia with an Atypical Presentation: Paroxysmal Hypoventilation Attacks**  
Ozturk Z., HIRFANOĞLU T., İNCİ A., OKUR İ., KOÇ E., TÜMER L., ARHAN E., Aydın K., SERDAROĞLU A.  
JOURNAL OF PEDIATRIC NEUROSCIENCES, cilt.13, sa.2, ss.276-278, 2018 (ESCI İndekslerine Giren Dergi)

## Kitap & Kitap Bölümleri

- I. **Yağ Asidi Oksidasyon Bozuklukları**  
İNCİ A., OKUR İ.  
TEMEL PEDIATRİ, HASANOĞLU ENVER, DÜŞÜNSEL RUHAN, BİDECİ AYSUN, BODUROĞLU KORAY, Editör, GÜNEŞ TIP KİTABEVLERİ, Ankara, ss.1126-1130, 2020
- II. **Normal Çocuklukta Beslenme ve Beslenme Bozuklukları**  
TÜMER L., İNCİ A., OKUR İ., Kasapkara Ç. S., OLGAÇ M. A. B.  
Lange - Current Tanı ve Tedavi Pediatri, Prof.Dr. Enver Hasanoğlu Prof.Dr. Aysun Bideci Prof.Dr. Elif N. Özmert Prof.Dr. Sevcan A. BAKKALOĞLU EZGÜ, Editör, ema tıp kitapevi, ss.281-308, 2018
- III. **Mitokondriyal Hastalıklar**  
TÜMER L., İNCİ A.  
Yurdakök Pediatri, Murat Yurdakök, Editör, Güneş Tıp Kitapevi, Ankara, ss.1779-1790, 2017
- IV. **Vitamin K**  
İNCİ A.  
Yurdakök Pediatri, Murat Yurdakök, Editör, Güneş Tıp Kitapevi, Ankara, ss.1552-1556, 2017

## Hakemli Kongre / Sempozyum Bildiri Kitaplarında Yer Alan Yayınlar

- I. **Screening of twelve lysosomal storage diseases with LC-MS/MS in Gazi university hospital in Turkey: The first results of validation**  
BİBEROĞLU G., İNCİ A., DERİN B., OKUR İ., EZGÜ F. S., TÜMER L.  
SSIEM, 3 - 06 Eylül 2019
- II. **Beneficial Effects of Modified Atkins Diet in Glycogen Storage Disease Type IIIa**  
OLGAÇ M. A. B., İNCİ A., OKUR İ., Kasapkara Ç. S., BİBEROĞLU G., OĞUZ A. D., Aktaş E., EZGÜ F. S., TÜMER L.  
SSIEM 2019, 3-6th September, 2019, Rotterdam-The Netherlands, 3 - 06 Eylül 2019
- III. **Next generation DNA sequencing as an initial diagnostic method for congenital defects of glycosylation**  
EZGÜ F. S., İNCİ A., Çiftçi B., TÜMER L., OKUR İ., Topçu B., Hasanoğlu A.  
SSIEM 2019, 3-6th September, 2019, Rotterdam-The Netherlands, 3 - 06 Eylül 2019
- IV. **Screening of Twelve Lysosomal Storage Diseases with LC-MS/MS in Gazi University Hospital: The First Results of Validation.**  
BİBEROĞLU G., İNCİ A., DERİN B., OKUR İ., EZGÜ F. S., TÜMER L.  
INTERNATIONAL INBORN ERRORS OF METABOLISM AND NUTRITION CONGRESS10 - 14 April 2019 Istanbul-Turkey, 10 - 14 Nisan 2019
- V. **Cornelia de Lange Syndrome and Glycogen Storage Disease Together in a Patient**  
KILIÇ A., EMECAN ŞANLI M., ÖZSAYDI E., İNCİ A., OKUR İ., TÜMER L., EZGÜ F. S.  
International Inborn Errors Of Metabolism And Nutrition Congress, İstanbul, Türkiye, 10 - 14 Nisan 2019
- VI. **A Very Rare Disease: Hyperornithinemia-Hyperammonemia-Homocitrullinuria (Hhh) Syndrome**  
Özsaydı E., Emecan Şanlı M., Kılıç m., İNCİ A., OKUR İ., TÜMER L., EZGÜ F. S.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- VII. **Novel Mutation in FBP1 Gene Presenting with Recurrent Episodes of Vomiting in A Child**  
Emecan Şanlı M., Kılıç m., Özsaydı E., İNCİ A., OKUR İ., EZGÜ F. S., TÜMER L.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- VIII. **Familial Hyperphosphatemic Tumoral Calcinosis in an Unusual Site**  
Emecan Şanlı M., Özsaydı E., Kılıç m., İNCİ A., OKUR İ., EZGÜ F. S., TÜMER L.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019

- IX. Hyperinsulinemic Hypoglycemia: Think of GLUD1 Gene Mutation Leading To Hyperinsulinism/Hyperammonemia (HI/HA) Syndrome**  
Emecan Şanlı M., Kılıç m., Özsaydı E., İNCİ A., OKUR İ., TÜMER L., EZGÜ F. S.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- X. Novel Mutation in Two Siblings with Normouricemic Lesch Nyhan Syndrome**  
Emecan Şanlı M., Özsaydı E., Kılıç m., İNCİ A., OKUR İ., EZGÜ F. S., TÜMER L.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- XI. A Novel Rars2 Mutation in Two Siblings with Microcephaly, Seizures and Liver Involvement**  
EMİNOĞLU F. T., Sevinç S., Karaköse Gök T., EZGÜ F. S., İNCİ A., TÜMER L.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- XII. Growth Hormone Treatment: Reverses Catabolic Process in Inborn Errors of Metabolism**  
İNCİ A., OKUR İ., AKKUZU E., DÖĞER E., BİBEROĞLU G., KALKAN G., TÜMER L., EZGÜ F. S.  
International Inborn Errors Of Metabolism And Nutrition Congress 10 - 14 April 2019, Istanbul-Turkey, 10 - 14 Nisan 2019
- XIII. Could Targeted Next Generation Sequencing Be A First Line Diagnostic Method for Lysosomal storage Diseases**  
İNCİ A., OKUR İ., AKKUZU E., DÖĞER E., BİBEROĞLU G., KALKAN G., TÜMER L., EZGÜ F. S.  
INTERNATIONAL INBORN ERRORS OF METABOLISM AND NUTRITION CONGRESS 10 - 14 April 2019 Istanbul-Turkey, 10 - 14 Nisan 2019
- XIV. RAR2 mutation in two siblings with microcephaly, seizures and liver involvement**  
EMİNOĞLU F. T., s s., gök t., EZGÜ F. S., İNCİ A., TÜMER L.  
15 th MEMG, Beyrut, Lübnan, 29 Kasım - 02 Aralık 2018
- XV. Respiratory system involvement of 41 Mucopolysaccharidosis patients with the evaluation of KL-6, SPA and SPD levels**  
İNCİ A., OKUR İ., Yılmaz Demirtaş C., BİBEROĞLU G., ASLAN A. T., EZGÜ F. S., TÜMER L.  
15 th MEMG, Beyrut, 29 Kasım - 02 Aralık 2018
- XVI. UNIQUE CLINICAL AND MOLECULAR FINDINGS IN LARGE COHORT OF PATIENTS WITH GAUCHER DISEASE FROM TURKEY**  
Akay Tayfun G., OKUR İ., BİBEROĞLU G., TÜMER L., İNCİ A., Küçükcongür A., Hasanoğlu A., EZGÜ F. S.  
Gaucher Symposium, İstanbul, Türkiye, 21 - 22 Ekim 2018
- XVII. Gaucher Disease and Pregnancy**  
İNCİ A.  
Gaucher Symposium, İstanbul, Türkiye, 21 - 22 Ekim 2018
- XVIII. Respiratory system involvement of mucopolysaccharidosis patients with the evaluation of KL-6, SPA and SPD levels**  
İNCİ A., OKUR İ., YILMAZ-DEMİRTAŞ C., BİBEROĞLU G., aslan a., EZGÜ F. S., TÜMER L.  
SSIEM, 4 - 07 Eylül 2018
- XIX. The clinical evaluation of Fabry patients with Mainz severity score index and DS3 score**  
OKUR İ., İNCİ A., bütün s., BİBEROĞLU G., EZGÜ F. S., TÜMER L.  
SSIEM, 4 - 07 Eylül 2018
- XX. Determination of succinylacetone in dried blood spot: preliminary results of our laboratory**  
BİBEROĞLU G., TÜMER L., OKUR İ., EZGÜ F. S., İNCİ A.  
SSIEM, 4 - 07 Eylül 2018
- XXI. Glycogen storage disease type 9: Insidious onset, mild form**  
TÜMER L., İNCİ A., OKUR İ., BİBEROĞLU G., EZGÜ F. S.  
SSIEM, 4 - 07 Eylül 2018
- XXII. An early diagnosis cerebretendinous xanthomatosis in a patient at the age of 15 years**  
İNCİ A., BİBEROĞLU G., OKUR İ., TÜMER L., EZGÜ F. S.

SSIEM, 4 - 07 Eylül 2018

- XXIII. **RENAL INVOLMENT IN FABRY DİSEASE**  
İNCİ A., BİBEROĞLU G., PAŞAOĞLU Ö. T. , TÜMER L., PAŞAOĞLU H., EZGÜ F. S.  
14 th middle east metabolic group (MEMG) meeting Athens GREECE, Atina, Yunanistan, 9 - 11 Şubat 2018
- XXIV. **Carnitine Acyl Carnitine TranslocaseDeficiency With Severe Hyperammonemiaand Hypoglycemia**  
İNCİ A., OKUR İ., OLGAC M. A. B. , AKKUZU E., BİBEROĞLU G., EZGÜ F. S. , TÜMER L.  
ICIEM, 5 - 08 Eylül 2017
- XXV. **Renal Involvement in Fabry Disease**  
İNCİ A., BİBEROĞLU G., OKUR İ., PAŞAOĞLU Ö. T. , TÜMER L., PAŞAOĞLU H., EZGÜ F. S.  
ICIEM, 5 - 08 Eylül 2017
- XXVI. **Preliminary Results of Our Laboratoryfor Bile Acid Metabolism Disorders**  
BİBEROĞLU G., DERİN B., İNCİ A., OKUR İ., EZGÜ F. S. , TÜMER L.  
ICIEM, 5 - 08 Eylül 2017
- XXVII. **Short Chain Fatty Acid OxidationDefect in an Adult Patient With RefractorySeizures**  
İNCİ A., TÜMER L., OKUR İ., BİBEROĞLU G., EZGÜ F. S.  
ICIEM, 5 - 08 Eylül 2017
- XXVIII. **Screening ALPL Gene Differences byNext Generation Sequence Techonology inPatients Having Low ALP Levels**  
İNCİ A., EZGÜ F. S. , topcu b., çiftci b., OKUR İ., BİBEROĞLU G., TÜMER L.  
ICIEM, 5 - 08 Eylül 2017
- XXIX. **DiagnosticCapability ofNextGenerationDNA Sequencing With A 450 Gene Panel forInborn Errors of Metabolism**  
EZGÜ F. S. , çiftci b., topcu b., İNCİ A., OKUR İ., BİBEROĞLU G., hasanoğlu a.  
ICIEM, 5 - 08 Eylül 2017
- XXX. **Renal İnvolvement in Fabry Disease**  
İNCİ A., BİBEROĞLU G., OKUR İ., PAŞAOĞLU Ö. T. , TÜMER L., PAŞAOĞLU H., EZGÜ F. S.  
13.International Congress of Inborn Errors of Metabolism., Rio de Janeiro, Brezilya, 5 - 08 Eylül 2017
- XXXI. **İnvestigation of LDLR Gene Mutations in Turkish Patients With Familial Hypercholesterolemia**  
OKUR İ., İNCİ A., OLGAC M. A. B. , ÇİFTÇİ B., TOPÇU B., TÜMER L., EZGÜ F. S.  
13th International Congress of Inborn Errors of Metabolism - ICIEM 2017, 5 - 08 Eylül 2017, cilt.5
- XXXII. **Ailevi Hiperkolesterolemi Olan Türk Hastalarda LDLR Gen Mutasyonlarının Araştırılması**  
OKUR İ., EZGÜ F. S. , İNCİ A., OLGAC M. A. B. , TÜMER L.  
2. Ege Endokrin Hastalıklar ve Genetik Sempozyumu, Türkiye, 23 - 25 Şubat 2017
- XXXIII. **Evaluation of chitotriosidase and high sensitive c reactive protein levels in mucopolysaccharidosis**  
İNCİ A., GENÇ B., YILMAZ-DEMİRTAŞ C., UDGU B., KARAOĞLU A., OKUR İ., EZGÜ F. S. , BİBEROĞLU G., TÜMER L.  
13th Middle East Metabolic Group Meeting/ Amman-Jordan, 28 - 30 Ekim 2016
- XXXIV. **Evaluation of chitotriosidase and high sensitivity c reactive protein levels in mucopolysaccaridosis patients**  
İNCİ A., DERİN B., YILMAZ C., udgu b., KARAOĞLU A., OKUR İ., EZGÜ F. S. , BİBEROĞLU G., TÜMER L.  
MEMG, 28 - 30 Ekim 2016
- XXXV. **Evaluation of chitotriosidase and high sensitive c reactive protein levels in mucopolysaccaridosis**  
İNCİ A., GENÇ B., YILMAZ-DEMİRTAŞ C., UDGU B., KARAOĞLU A., OKUR İ., EZGÜ F. S. , BİBEROĞLU G., TÜMER L.  
13th MEMG Meeting, 28 ekim-30kasım 2016, Amman, Jordan, 28 - 30 Ekim 2016
- XXXVI. **Could propionylcarnitine and free carnitinebe used as antioxidative markers in mucopolysaccaridosis**  
İNCİ A., BİBEROĞLU G., DERİN B., KARAOĞLU A., OKUR İ., EZGÜ F. S. , TÜMER L.  
MEMG, 28 - 30 Ekim 2016
- XXXVII. **Do cytokine levels play a role in the pathogenesis of mucopolysaccharidosis patients**  
İNCİ A., TÜMER L., YILMAZ-DEMİRTAŞ C., KARAOĞLU A., OKUR İ., OLGAC M. A. B. , EZGÜ F. S. , BİBEROĞLU G.  
13th Middle East Metabolic Group Meeting/Amman -Jordan, 28 - 30 Ekim 2016
- XXXVIII. **Early initiation of investigational enzyme replacement therapy in a nine month old infant with**

**mucopolysaccharidosis type VII**

KARAOĞLU A., İNCİ A., BİBEROĞLU G., OKUR İ., Kılıçkaya a., TÜMER L., King b., haller c., EZGÜ F. S.  
MEMG, 28 - 30 Ekim 2016

- XXXIX. **Evaluation of chitotriosidase and high sensitivity c reactive protein levels in mucopolysaccharidosis**  
İNCİ A., Genç B., Demirtaş C., Udgu B., KARAOĞLU A., OKUR İ., EZGÜ F. S. , BİBEROĞLU G., TÜMER L.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XL. **Early initiation of investigational enzyme replacement therapy in a 9 month old infant with mucopolysaccharidosis type VII**  
KARAOĞLU A., İNCİ A., BİBEROĞLU G., OKUR İ., Kılıçkaya A., KELEŞ E., TÜMER L., King B., Hall C., EZGÜ F. S.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLII. **Bone mineral density and vitamin D status in inborn errors of metabolism**  
OLGAÇ M. A. B. , TÜMER L., İNCİ A., KARAOĞLU A., OKUR İ., EZGÜ F. S.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLIII. **Do cytokine levels play a role in pathogenesis of mucopolysaccharidosis patients**  
İNCİ A., TÜMER L., Demirtaş C., KARAOĞLU A., OKUR İ., OLGAÇ M. A. B. , EZGÜ F. S. , BİBEROĞLU G.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLIV. **Identification of a novel mutation in Turkish infant with early onset monocarboxylate transporter 1 MCT1 deficiency as a cause of recurrent ketoacidosis**  
OKUR İ., İNCİ A., KELEŞ E., KARAOĞLU A., Ceylaner S., BİBEROĞLU G., EZGÜ F. S. , TÜMER L.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLV. **Type 1 hypersensitivity reaction and desensitization with Elosulphase alpha**  
İNCİ A., Kan A., Topuz B., OKUR İ., EZGÜ F. S. , TÜMER L.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLVI. **BonemineradensityandvitaminDstatusininbornerrorsofmetabolism**  
OLGAÇ M. A. B. , TÜMER L., İNCİ A., KARAOĞLU B., OKUR İ., EZGÜ F. S.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016
- XLVII. **Identification of a novel mutation in Turkish infant with early onset monocarboxylate transporter 1 MCT1 deficiencyasacauseofrecurrent ketoacidosi**  
OKUR İ., İNCİ A., KELEŞ E., KARAOĞLU A., CEYLANER S., BİBEROĞLU G., EZGÜ F. S. , TÜMER L.  
SSIEM 2016: Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Rome, Italy, 6 - 09 Eylül 2016, cilt.39, ss.35-284
- XLVIII. **Plasma acylcarnitine levels Are there New İnflammatory markers in lysosomal storage disease**  
BİBEROĞLU G., DERİN B., İNCİ A., udgu b., kurnaz p., OKUR İ., EZGÜ F. S. , TÜMER L.  
MEMG, 29 Ekim - 01 Kasım 2015
- XLIX. **Is there any effect of acylcarnitines on proinflammatory process in obese children**  
BİBEROĞLU G., DERİN B., İNCİ A., DÖĞER E., OKUR İ., EZGÜ F. S. , TÜMER L.  
SSIEM, 1 - 04 Eylül 2015
- L. **Patient with Niemann Pick type C presenting with lymphatic involvement with Niemann Pick cells in the left jaw**  
İNCİ A., OKUR İ., ESENDAĞLI G., OKUR A., EZGÜ F. S. , TÜMER L.  
Annual Symposium of the Society for the Study of Inborn Errors of Metabolism, Lyon, Fransa, 1 - 04 Eylül 2015
- L. **Mucopolysaccharidosis Type VII at an Early Age A good candidate for investigational enzyme replacement therapy**  
Abdulkaki K., EZGÜ F. S. , BİBEROĞLU G., OLGAÇ M. A. B. , İNCİ A., TÜMER L.

SSIEM, 1 - 04 Eylül 2015

- LI. A completely new approach to the diagnosis of inborn errors development of a 450 gene all metabolic disorders next generation sequencing panel**  
EZGÜ F. S. , çiftçi b., topçu b., OKUR İ., İNCİ A., OLGAÇ M. A. B. , KARAOĞLU A., BİBEROĞLU G., TÜMER L., hasanoğlu a.  
SSIEM Annual Symposium, 1 - 04 Eylül 2015
- LII. Dihydrolipoamide dehydrogenase deficiency diagnosed by using new generation sequencing technology**  
İNCİ A., TÜMER L., OKUR İ., OLGAÇ M. A. B. , SARI S., çiftçi b., topçu b., EZGÜ F. S.  
SSIEM Annual Symposium, 1 - 04 Eylül 2015
- LIII. Sol çenede Lenfatik tutulum ile giden Niemann Pick tip C olgusu**  
İNCİ A., OKUR İ., ESENDAĞLI G., OKUR A., OLGAÇ M. A. B. , EZGÜ F. S. , TÜMER L.  
XIII.Ulusal Metabolik Hastalıklar ve Beslenme Kongresi, Adana, Türkiye, 14 - 18 Nisan 2015

## **Atıflar**

Toplam Atıf Sayısı (WOS):3

h-indeksi (WOS):1